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AN INTRODUCTION

TO THE

CHEMISTRY OF PLANT PRODUCTS

BY

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PREFACE TO THE SECOND EDITION.

The great advances made in the chemistry of Plant Pigments since the issue of the first edition have necessitated the re-writing of the section dealing with this subject. For the rest, we have confined ourselves to making a few minor additions and corrections and adding a number of further references to the literature.

P. H. T. G. H.

July, 1916.

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PREFACE TO THE FIRST EDITION

The importance to the botanist of a working knowledge of chemistry can hardly be overestimated, since vegetable physiology is replete with problems awaiting solution by the combined application of botanical and chemical methods.

Teachers of vegetable physiology, however, not infrequently find that their students' knowledge is deficient in just those branches of chemistry which are of particular importance to the botanist, which is, no doubt, largely due to the fact that those compounds which are of interest to the botanist do not necessarily fit into the scheme of instruction of the chemist.

The present work is an attempt to provide such students, who are assumed to have some acquaintance with chemistry, with an introductory account of the chemistry and biological significance of some of the more important substances occurring in plants.

In compiling this book various sources of information have been laid under contribution, and although the point of view is, in the main, purely chemical and botanical, the economic aspect has not been lost sight of, and, where possible, mention has been made of the uses of plant products and of the manufacturing processes employed in their preparation.

P. H. T. G. H.

December, 1912.

CONTENTS.

									P	AGE
PREFACE										V
SECTION I FATS, OILS,	AND	WA	XES	. 1	PHOS	PHA	TIDI	ΞS		I
Fats		_								1
Occurrence .				Ċ		Ċ	:		:	ī
Industrial uses of	veget	able	fats a	und :	oils					3
Constitution .										5
										10
Extraction . Physical properties	es									11
Chemical properti	es				:					12
Saponification										13
Saponification Cholesterol and Phyto Spontaneous changes	sterol									15
Spontaneous changes	in fat	8								18
Rancidity .										18
Drying and resinit	icatio	n						÷	Ċ	19
General properties and	react	tions	of fa	te.	•	•	Ċ		·	20
Special tests for partic	nlar c	lasse	s of f	ate	•	•	•		٠	21
Special tests for partic Colour reactions . Microchemical reactio		111000			•	•	•	•	•	22
Microchemical reaction	ns	•	•	•	•	•	•	:	•	23
Quantitative estimation	1 .				÷			•	:	24
Quantitative methods			racte	riza	tion c	of fats	•	•	•	28
Physiological significa	nce						•		·	35
Waxes		•	•	•	•			:		
Phosphatides, Lecithins, or	Dhou	· nhali	· nince					•		43
Occurrence	FIIOS	phon	pines	٠.	•	•	•	•	•	4.1
Occurrence Preparation	•	•	•	•	:	•	•	•	•	44
Reactions and characte	omiotia	•	•	•	•	•	•	•	•	45
Choline	cristic	8	•	•	•	•	•	٠	•	46
Choline Formation of Lecithin	•	•	•	•	•	•	•	•	•	47
Physiological significa		•	:	:		•	•	•	•	49
i nysiologicai signinea	lice	•	•	•	•	•	•	•	٠	50
anamian II albhailte	m .	3.63								
SECTION II.—CARBOHYD			•	٠	•	•	٠	•	•	52
Classification				•						52
Constitution and isomerism	of su	igars			•		•			5 +
General reactions of sugars			٠	•		•				56
Monosaccharides		•	•	٠	•	•				57
Pentoses General properties			•		•					57
General properties	s .									57
Properties of indi-	vidual	pent	oses			:				59
Hexoses										60
Glucose or dextrose										60
Preparation .										60
•		vii							•	

CONTENTS

1)										P 2	rer.
Properties Reactions Microchemica Levulose or Fruct Properties Reactions Sorbose Galactose Preparation Properties Detection Mannose Detection Disaccharides Cane sugar, sucro Properties and Maltose Properties and Isomaltose Cellobiose Mycose or trehalo Agavose and lupee Lactose or milk st Trisaccharides Raffinose Detection Melecitose Sugars of unknown me Estimation of sugars Volumetric metho			•	•		•	•				61
Keactions	:		•	•	•	•	•				61
Microchemica	l test	S		•	•	•					63
Levulose or Fruct	ose		•				•	•			64
Properties	•	•		•		•	•				65
Reactions	•	•	•			•	•				65
Sorbose .	•	•		•		•	•	•	•		65
Galactose .	•	•					•				65
Preparation		•									66
Properties	•			•			•	•			66
Detection		•				•					66
Mannose .	•	•				•		•			67
Detection	•						•				67
Disaccharides .							•				68
Cane sugar, sucros	se or	sacci	arose	3		•	•				68
Properties and	d reac	ctions									70
Maltose .	•										71
Properties and	d reac	ctions	,								72
Isomaltose .											73
Cellobiose .											73
Mycose or trehalo	se										73
Agavose and lupe	ose										73
Lactose or milk su	ıgar										74
Trisaccharides .											74
Raffinose .											74
Detection											75
Melecitose .											76
Sugars of unknown me	olecul	ar w	eight	or su	gar-li	ke po	lvsac	chari	les		76
Estimation of sugars											77
Volumetric metho	ds										77
Estimation of sugars Volumetric metho Estimation by Estimatic Estimatic Estimatic Estimatic Estimatic Estimatic Estimation Estimation by Gravimetric metho	z mea	ns of	Febl	inσ's	solut	ion				•	77
Estimation	on of	nento	nses					•	•	•	78
Estimatio	on of	oluce	nse.	•	•	•	•	•	•	•	70
Estimatio	on of	galac	tose	ond r	· nann	•	•	•	•	•	/9
Estimatio	on of	cane	211030			J	•	•	•	•	8.
Estimatio	on of	malt	ougai		•		•	•	•	•	01
Estimatio	on of	mint	J8C	fana	•	•	•	•	•	•	01
Estimation be)11 ()1	mixtu	Dane	l sug	a15	•	•	•	•	•	02
Estimation by	. Don	uis oi	ravy	5 50	iutioi		•	•	•	•	03
Estimation by Estimation by Gravimetric methe Estimatic Estimatic Estimatic Polarimetric meth Polysaccharides . Classification	/ Den	leaict	s son	HOIL	•	•	•	•	•	•	05
Gravimetric metric	oas			•	•	•	•	•	•	•	80
Estimatio	on or	penti	oses	•	•	•	•		•	•	00
Estimatio	on or	giuco	se	•	•	•	•	•	•	•	88
Estimatio	on or	gruce	ise as	osaz	one	•	•	•	•	•	92
Polarimetric meth	ods	•	•	•	•	•	•	•	•		93
Polysaccharides .	•	•	•	•	•	•	•	•	•	•	96
Classification	•	•	•	•	•	•	•	•		٠	96
Starches .	:	•	•	•	•	•	•			•	97
Starch or am	ylum	•	•	•	•	•		•	•	•	97
Preparati	ion		•	•						٠	98
Propertie	es								•	•	99
Action of	t acid	s on	starch	1	•					•	102
Polarimetric meth Polysaccharides . Classification Starches . Starch or am Preparati Propertic Action o Action o Action o Reaction Estimatic	f mal	t or d	iastas	e on	starc	h		•	•		103
Action of	f bact	eria o	on sta	rch						٠	103
Reaction	IS									٠	104
Estimation	on										105

											PAGE
. Dextrins											TOD
Genera	l prope	erties			•	•	•	•	•	•	100
Amylor	levtrin		•	•	•	•			•		109
Frythre	devtri		•	•		•				•	109
Commo	roiol d	10			•				•		110
Dextrins Genera Amyloc Erythre Comme Glycogen Prepare Propert Identifi Estima	iciai c	extrin			•		•		•		110
Glycogen	.:								•	•	110
Prepara	ition		•								112
Propert	ies										113
Identifi	cation										114
Estima	tion										114
Paradextran	e and	paraiso	odext	rane.							114
Estima Paradextran Inulin . Prepara											114 114 114
Prepara	ition .							•		•	TT7
Charact Identific Inulin-like s	erc	•	•						•	•	117 118 118
Identific	cation		•						•		0
Invlin like a	uboton								•		110
Manual S	uostan	ces .	•					•			I20 I20
Mannane	•		•						•	•	120
Paramannar	ie										121
Carubin											121
Galactane											122
Inulin-like s Mannane Paramannar Carubin Galactane Paragalacta Amyloid Gums . Microchemic Gum tragac Wood gum Wound gum Wound gum Mucilage Functio Pectic bodie: Microch Cellulose . Classification Characterist Action of ch Characterist Constitution Industrial us	ne .										122
Amyloid											122
Gums							•		•	•	122
Microchemic	al read	tions	•		•	•	•	٠	•	•	123
Gum arabic	ai ica	cions	•				•	•	•	•	124
Ponetio	• •	•	•	•		•	•				125
Reactio	ns .		•	•	•	•	•		•		125
Gum tragac	anth .	•	. •								126
Wood gum	and ce	rasın c	or che	erry g	um .						126
Wound gum	١.										126
Mucilage											127
Functio	n.										128
Pectic bodies	S .										128
Microch	emical	reacti	ons						Ĭ.		130
Cellulose .								•	•		727
Classification	, .	•	•	•	•	•	•	•	•	•	131
Characterist	ice and		rtion.	of no	1	aclini	•	•	•	•	131
Antion of al	cs and	prope	111.	01 110	rmai	cenui	ose	•	•	•	133
Action of ch	emicai	s on ce	iluio	se .	•			•	•	•	134
Characteristi	cs and	prope	rties	of co.	mpcu	nd cel	lulose	s .			137
Constitution Industrial us	٠.,٠	· •	•								139
											141
Commerciall	y valu	able de	erivat	ives o	of cel	lulose		-			141
Microchemic	al reac	tions									144
The synthesis of	carboh	ydrate	es in	green	plan	ıts .					146
Introductory	: Alde	hydes			٠.						146
Formale	lehvde	٠.									151
Photosynthe	sis .					•	•	•	•	•	75.
Microchemic The synthesis of Introductory Formale Photosynthe	•	•		•	•	•	•	•	•	•	154
SECTION III.—GL	ucos	IDES									-6-
Constitution	200.,	2010	•	•	•	•		•	•	•	169
Identification		•	•	•	•	•					
Dhamister				•		•		•	•		
Physiological sig	шпсан	ce .			•			٠			173
Cyanogenetic gli	1coside	es .									176
Isolation		-									178
Chemistry											178
Constitution Identification Physiological sig Cyanogenetic glt Isolation Chemistry Reactions											179

											р	AGE
Amygdalin Dhurrin Phascolunat Lotusin. Saponins General pro Solubility Physiologice Chemistry Reactions Other glucosides Sinigrin											-	181
Dhurrin	•	•	•	•	•	•	•	•	•			182
Phaseolunat	in.	•	•	•	•	•	•	•		:		182
Lotuein	•••	•		•	•	•	•	•			:	
Saponins .	•	•	•	•	•	•	•	•			:	183
General pro	nertie	· · and		•	•	•	•				:	~~
Solubility	pertie	and	uses	•	•	•		•				184
Physiologics	Lacti	•	•	•	•	•	•	•	•		:	185
Chemistry	ii actii	UII		•		•		•				185
Posstion	•	•	•	•	•	•			:		:	187
Reactions Other glucosides	•	•	•	•	•	•	•	•			•	187
Other glucosides	•	•	•	•	•		•			•	•	187
Sinigrin Prepara	.:	•	•		•	•	•			•	٠	187
								•	•	•	•	
Coniferin Salicin . Prepara	•	•		•	•	•	•	•	•		•	188
Salicin .	.:	•		•	•	•	•	•			•	
Prepara	tion		•		•	•	•				•	189
											٠	191
Identific	cation					•						191
SECTION IV.—TA	NNIN	IS	•					•		•	٠	193
Occurrence .								•	•			194
Microchemical re	eaction	ns										197
Chemistry .												200
Microchemical re Chemistry . Pyrocatechol, ca	techol	, or p	угоса	atechi	n							201
Pyrocatechol, ca Reactions Resorcinol . Reactions Hydroquinone Reactions Protocatechuic a Reactions Pyrogallol or pyrogallol or pyrogallol												202
Resorcinol .												202
Reactions												202
Hydroguinone						:						202
Reactions											Ċ	202
Protocatechnic a	cid											203
Reactions			•	•		•	•	•				203
Purogallol or py	rogall	ic aci	A	•	•				•	•	:	204
Reactions	ogan	ic aci	u	•	•				•			204
TOLI 1 1 1	•			•	•		•		•		٠	
Parations	•	•	•	•	•	•	•			•	•	204
Priorogiucinol Reactions Gallic acid . Reactions Ellagic acid . Reactions Philobaphenes Tannins as glucc Classification of Properties and d Pyrogallol t	•	•	•	•	•	•	•	•	•		٠	205
Gaine acid .	•	•	•	•	٠	•	•	•	•		•	
Reactions	•	•	•	•			•	•		•	•	206
Ellagic acid.		•	•	•		•	•	•		•		206
Reactions	•	•	•	•	•	•	•	•			٠	207
Phlobaphenes			•		•							208
Tannins as gluce	sides											208
Classification of	tanni	ns										209
Properties and d	escrip	tion (of ind	lividu	al tar	nins						211
Pyrogallol t	annins	š										211
Gallota	nnic a	cid										211
Ex	tractio	n										213
Re	action	S										214
Pyrogallol t Gallota Ex Re De	tection	n of g	allic	acid	in pre	esence	of g	allota	nnic a	acid		214
Co	nstitut	ion										214
Co Ellagita	nnic	acid									:	216
Pyrocatecho Catechi	l tann	ins	-						•		:	216
Catachi	ı tanı	ic aci	id.	•	•		:	•	•	•		216
('atachi	n tailli	ic at		•	•	•	•	•	•	•	٠	
Catechi Quercit Physiological sig	u annia	· acid	•	•	•	•	•	•	:	•	•	217
Physiological -:-	anne milier	acid	f ton	·	•	•	•	:	•	•	•	
r nysiological sig	пппса	nice 0	ı tanı	ums								217

CONTENTS

хi

SECTION V.—PIGMEN' Chlorophyll Constitution . Action of alkalies Action of acid Crystalline and an Relationship betw Extraction of chlo Carotinoids or yellow Carotin . Yantbonhyll	mc									F	PAGE
SECTION V.—PIGMEN	15	•	•	•	•	•	•	•	•	٠	225
Chlorophyll	•	•	•	•	•	•	•	•		•	225
Constitution .	•	•	•	٠	•	•	•	•		•	231
Action of alkalies	٠	•		•	•	•	•	•	•	•	232
Action of acid	٠.			٠.		•	•	•	•		233
Crystalline and an	norp.	hous	chloro	phy	II .	٠.		•	•	•	233
Relationship betw	een.	chlor	ophyll	an	d hæi	nogle	obin		•	٠	230
Extraction of chlo	roph	yll	•	•		.:.	;		٠	٠	230
Carotinoids or yellow	pigir	ents	accon	ipar	yıng	chlor	opny	11 .	•	•	241
Carotin	•	•		•	٠	٠		•	•		242
Xanthophyll .	•	•	•	•	•	•	•	•	•	•	243
Fucoxanthin.	•		•	•	•	•	•	•	•	•	243
Anthoxanthins .	•	•	•	•	•	•		•		•	244
Flavones and Xanthon	ı€S	•	٠	•	•	•		•	•	•	244
Yellow colouring	mati	ers d	lerived	fro	n flav	one	•	•	•	•	245
Yellow colouring	matt	ers d	lerived	fro	m xar	thon	е.	•			246
Properties of anthoxar	nthin	IS						•	•		247
Anthocyanins, Phycoe	rythi	rin, a	nd Ph	yco	haeir	· .	•				248
Connection between a	ntho	cyan	ins and	i an	thoxa	nthir	ns .				251
Extraction of anthocy	anin:	s.									251
Anthocyanins .											252
Properties .											254
Reactions .											256
Physiological sign	ifica	nce									256
Phycoerythrin .											258
Preparation .											259
Reactions .											259
Carotin . Xanthophyll . Fucoxanthin . Anthoxanthins Flavones and Xanthon Yellow colouring Yellow colouring Properties of anthoxan Anthocyanins, Phycoc Connection between a Extraction of anthocy Anthocyanins Properties Reactions Physiological sign Phycocytythrin Preparation Reactions Phycophaein				٠						•	26 0
SECTION VINITRO	1EN	BA	SES								263
Aikaloids				•	•	•		·	Ĭ.		265
Classification		•	•		•	•		Ċ	·		256
Properties	•	•	•	•	•	•	•	•	·		260
Reactions .	•	•	•	•	•	•	•	•	•	•	270
Isolation .	•	•	•	•	•	•	•	•	•	•	271
Origin of alkaloid	e in	the r	dent.	•	•	•	•	•	•	•	272
Ptomaines		tile j	rianic	•	•	•	•	•	•	•	274
Purine bases	•	•	•	•	•	•	•	•	•	•	276
Physiological significa		of n	tracen		•	•	•	•	•	•	280
SECTION VI.—NITROC Alkaloids	ance	01 111	trogen	Da	cs	•	•	•	•	•	200
SECTION VII.—COLLO	oids		•		٠		•	•	•	•	283
Properties		•							•		284
Diffusibility .											284
Optical properties											285
Change of state o	r gel	forn	nation								287
Protective power	•										291
Classification of Collo	ids	. :									291
Properties of susp	ensc	oids									292
Properties of emu	ilsoid	ls									293
The nature of gel	İs										294
SECTION VII.—COLLO Properties Diffusibility . Optical properties Change of state o Protective power Classification of Collo Properties of susp Properties of emu The nature of gel Adsorption .											295

									P	AGE
SECTION VIII.—PROTEINS	. ·									301
General properties .										302
Microchemical reactions										308
SECTION VIII.—PROTEINS General properties Microchemical reactions Proteins as colloids										308
Microchemical reactions Proteins as colloids Decomposition products Amino acids obtained as cl										313
Amino acids obtained as cl	eavag	e pro	ducts	of pr	otein	S				315
Constitution of the protein	molec	cule								318
Constitution of the protein Polypeptides Occurrence of amino acids Classification of proteins Comparison between veget										319
Occurrence of amino acids	in pla	ints								321
Classification of proteins										322
Comparison between veget	able a	ınd ar	imal	prote	ins					328
Extraction of proteins .										330
Synthesis of proteins in the	: plan	t								333
Synthesis of amino acids in	n the	plant								337
Extraction of proteins . Synthesis of proteins in the Synthesis of amino acids in Estimation of nitrogen										340
2 minution of managem	•	•	•	•						٥.
SECTION IX.—ENZYMES										344
Classification	-									348
Isolation		Ċ				Ċ	Ċ			349
Chemical constitution .	•	:				Ċ	Ċ			349
Properties	•		•				:	Ċ		350
Colloidal nature of enzyme	· ·	•						Ĭ.		351
Mode of action		•	•	•		•	•			352
Activators	•	•	•	•	•	•		•		354
Paralyeers	•	•	•	•	•	•	•	:	:	
Anti-enzymes	•	•	•	•	•	•	·	•		358
Engrand the laws of		ction	•	•	•	:		•		361
SECTION IX.—ENZYMES Classification Isolation Chemical constitution Properties Colloidal nature of enzyme Mode of action Activators Paralysers Anti-enzymes Enzymes and the laws of r Consideration of certain ty Lipase Isolation Quantitative deter Diastase	nec of	Fenas	mec	•	•	:	:		•	365
Tiposo	pes or	CHZ	nics	•	•		•	•	•	366
Lipase	•	•	•	•	•		•	•		367
Oversitation deter	· :		· :		•	•	•	•		
Quantitative deter Diastase Isolation .	mma	tion o	acti	vity	•	٠	٠	•	٠	
Diastase	•	•	•	•	•		•	•		368
Isolation	٠.		·	•	•	•	•	•		369
Quantitative dete	rmına	tion c	ı actı	vity		٠	•	•		370
Proteases Isolation .	•	•	•	•	•	•	•	•		370
isolation .	:	•	•	٠		٠	•	•	•	٠.
General considera	itions	•	•		•	•	•	•		372
Tryptophane reac Quantitative dete	tion		•	•				:		373
Quantitative dete	rmina	tion c	of acti	vity						374
Zymase and alcoholic Isolation of zyma General considers Identification of e Quantitative dete Occurrence of alc	term	entati	on							374
Isolation of ∠yma	se									377
General considera	itions									377
Identification of e	thyl a	alcoho	ol							382
Quantitative dete	rmina	tion (of act	ivity						383
Occurrence of alc	ohols	in pla	ants							385
Inosite Manufacture of e								:		387
Manufacture of e	thyl a	lcoho	l.							389
										390
lsolation Peroxidase Preparation										392
Perexidase .								:		
Preparation										
Identification								:		393
General considers	ations									394
										031
INDEV										

SECTION I.

FATS, OILS, AND WAXES.

In ordinary parlance, no clear distinction is made in the use of the terms fat and wax, which are applied more or less indiscriminately to any solid substances which have a greasy feeling to the touch and do not mix with water. Chemically, however, there is a marked difference between the two classes; the fats are compounds of the trihydric alcohol glycerol, whereas the waxes are compounds of the higher monohydric alcohols, such as cetyl alcohol $C_{16}H_{33}OH$, myristic alcohol $C_{30}H_{31}OH$, and cholesterol $C_{27}H_{45}OH$.

The tendency to rely on physical properties only, and to regard waxes as having generally a harder consistency than fats has given rise to several cases of incorrect nomenclature. For example, wool fat and spermaceti being compounds of cholesterol and cetyl alcohol are in reality waxes, though they are usually regarded as fats, whereas the substance ordinarily known as Japan wax is actually a fat, since it is a compound

of glycerol.

The term oil, as used in the ordinary sense to imply a liquid which is immiscible with water, must not be taken to have any chemical significance, since substances having this physical property are found in almost every class of chemical compound. Used in connexion with fats, the term oil simply implies a fat that is liquid at ordinary temperatures; any solid fat on melting becomes an oil, and, on the other hand, any fatty oil on solidifying becomes a fat.

OCCURRENCE.

Fats are very widely distributed in the vegetable kingdom, the most common being the glycerides of oleic, palmitic, and stearic acids; especially are they found in reproductive bodies

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such as spores and seeds. Not only are the spores, both sexual and asexual of very many Algæ and the majority of the Fungi characterized by the presence of oil, but also the thallus: the filaments of *Vaucheria*, for instance, contain much oil; and of the Fungi, the sclerotia of *Claviceps purpurea* (ergot) may contain as much as 60 per cent, whilst the mycelium of *Lactarius deliciosus* contains about 6 per cent.

The fats of the Fungi are rich in fatty acids associated with lecithins and ergosterins.

In Angiosperms they are widely distributed, especially in seeds where they may replace the carbohydrates as a reserve food-material and are not uncommonly associated with protein reserves; to mention a few examples, colza oil is obtained from the seeds of Brassica Napus, palm oil from the fruits of Elaeis guineensis, cotton-seed oil from Gossypium herbaceum, linseed oil from Linum usitatissimum, olive oil from the sarcocarp of Olea europæa, castor oil from the seeds of Ricinus, and cacao butter from the fruits of Theobroma.

Oils of lesser economic importance occur in the fruits or seeds of the sunflower, almond, hemp, willow and many other plants.

The amount of oil present in such structures may be quite considerable, thus in the kernel of the Brazil nut nearly 70 per cent may obtain, and in the almond about 54 per cent.

Oils also occur in the vegetative organs to a greater or lesser extent; substances of an oily nature are found in association with the chloroplasts and, in some cases, to a relatively large extent, e.g. in *Strelitzia*; sometimes it is present as a definite reserve food-material as in the tubers of *Cyperus esculentus*, where it is associated with starch, and in the roots of some orchids.

This particular form of food reserve is doubly of value since its presence may lessen the danger arising from drought, and also more energy can be stored up in the form of oil than in an equal bulk of carbohydrate; in this connexion may be mentioned the fact that in some cases the appearance of oil may be transient, thus in some trees the starch stored up in the parenchyma of the stem may be converted into fat during the winter's cold; the starch, however, reappears on a rise in temperature. Also oil may appear in the leaves of evergreen

plants during the winter months. These phenomena are similar to those which will be mentioned in connexion with the conversion of starch into sugar under the influence of low temperature (p. 116).

The majority of vegetable fats are fluid at ordinary temperatures; a few, however, are solid, for instance, cacao butter and the fat in the seeds of *Myristica*.

INDUSTRIAL USES OF VEGETABLE FATS AND OILS.

Economically, fats are of considerable value, being used for food, illumination, lubrication, soap manufacture, and for a variety of other purposes.

The following is a brief consideration of some of the more important industrial uses of the commoner fats and oils of vegetable origin.

OLIVE OIL is extracted from the fleshy pericarp of the fruit of the olive, *Olea europaea*, by pressure. The best quality oil, which is expressed without the application of heat, is used for food; lower grade oils, obtained by extracting the residues from the presses with fat solvents, such as carbon disulphide or light petroleum, are used in the manufacture of soap (see p. 5).

COTTON-SEED OIL is extracted from the seeds of Gossypium herbaceum by pressing them at a temperature of about 90°; the crude brown oil is purified by treatment with caustic soda which removes the free fatty acids, colouring matter and other impurities. After purification the oil is light yellow in colour. It is used for the manufacture of soap and rubber substitutes.

Coco-Nut OIL is obtained from the ripe seeds of *Cocos nucifera* and *Cocos butyracea* by pressure; the dried endosperms, known as Copra, are imported into Europe and the oil extracted from them is commonly known as Copra oil. Soaps made from coco-nut oil have the property of absorbing large quantities of salt solutions and can, therefore, be used for washing with sea-water.

PALM OIL which occurs in the fruit of *Elaeis guineensis* is, when pure, a colourless substance of the consistency of lard; on exposure to air it readily turns yellow, but the colour can be removed by oxidation by means of a current of air. It has

a faint odour resembling that of violets. Both coco-nut oil and palm oil in the crude state contain free fatty acids which can, however, be removed by treatment with alcohol. When so purified they are employed as substitutes for butter under the name of vegetable butter or palmine, etc.

RAPE OIL or COLZA OIL is a thick, yellowish oil obtained from the seeds of *Brassica Rapa* and *Brassica Napus* which is used as an illuminant.

By drawing a current of air through the oil heated to 70° a so-called "blown" oil is produced, the specific gravity of which becomes almost equal to that of castor oil, namely 0.97; in this condition it is miscible with mineral oils. The mixture which is known as marine oil is used for lubricating marine engines.

LINSEED OIL is obtained by pressing the seeds of *Linum usitatissimum* either with or without the application of heat; the residues after compression are made up into cattle food.

The drying vegetable oils, particularly linseed oil, are used in the manufacture of oil paints as vehicles for the pigments; for artist's white paints, walnut and poppy-seed oils are chiefly used. The drying properties of linseed oil used for the manufacture of paint are greatly increased by boiling with lead oxide; such oil is known as boiled oil. A similar effect may be produced by dissolving in it certain salts known as "driers," such as lead linoleate or the metallic salts of resin acids, etc.

Varnish consists of a mixture of boiled oil with gum resins and oil of turpentine.

CASTOR OIL is obtained by compressing the seeds of *Ricinus communis* either with or without the application of heat. The seeds contain a fat-splitting enzyme which is employed commercially for the hydrolysis of fats; they also contain a very poisonous toxalbumin, known as Ricin, which remains in the residues after the expression of the oil. Castor oil is a thick viscid colourless liquid; when heated above 280° it decomposes with the formation of oenanthol, a substance having a very unpleasant odour. Castor oil is largely used in the dye industry; for this purpose it is converted into the so-called turkey red oil, used for alizarin dyeing, by treatment with sulphuric acid and neutralization of the resulting sulphonic acid with soda.

For the manufacture of soap the following fats and oils are used: tallow fat, palm oil, palm-kernel oil, coconut oil and olive oil; the fats are boiled with caustic soda until saponification is complete, whereupon the mixture is saturated with common salt. The soap, being insoluble in strong salt solution, rises to the surface leaving the glycerol and salt in the aqueous layer below; the latter is then run off and the scum, which is allowed to harden in moulds, is known as hard soap. Soft soaps are prepared by boiling the cheaper oils, such as hemp-seed oil, cotton-seed oil or linseed oil with caustic potash; when saponification is complete the mixture is allowed to set to a semi-solid without the addition of sodium chloride; the resulting mixture contains all the glycerol together with the excess of alkali and a quantity of water.

The whole of the glycerol of commerce is obtained from fats; it is used largely for the manufacture of dynamite.

CONSTITUTION OF FATS.

Fats belong to that class of organic compounds which are known as esters, an ester occupying in organic chemistry exactly the same position as salts do in inorganic chemistry.

When an inorganic base such as caustic soda or potash reacts with an acid, either organic or inorganic, one or more of the replaceable hydrogen atoms in the acid is replaced by the metal and the resulting product is known as a salt; thus:—

$$NaOH + H_2SO_4 = NaHSO_4 + H_2O$$

or $KOH + CH_3COOH = CH_3COOK + H_2O$
Acetic acid Potassium
acetate

If now the inorganic base be replaced by its organic analogue, an alcohol, a similar reaction ensues, but the resulting compound is called an *ethereal salt* or *ester*.

$$\begin{array}{cccc} C_2H_5OH + H_2SO_4 = C_2H_5HSO_4 + H_2O \\ & & Ethyl \ hydro-\\ gen \ sulphate \\ C_2H_5OH + CH_3COOH = CH_3COOC_2H_5 + H_2O \\ Ethyl & Acetic & Ethyl \ acetate \\ alcohol & acid & \end{array}$$

The reaction between an acid and an alcohol containing

three hydroxyl groups (OH) instead of only one, may be expressed by the following equation:—

the resulting compound tristearin also being an ester.

The naturally occurring fats are mixtures of similar esters of glycerol with other fatty acids such as palmitic $C_{15}H_{31}COOH$ or butyric C_3H_7COOH acids, or with the unsaturated acid oleic acid $C_{12}H_{33}COOH$.

A wax, on the other hand, is an ester of a monohydric alcohol as illustrated by the equation:—

$$\begin{array}{lll} C_{15}H_{31}COOH + & C_{30}H_{61}OH = & C_{15}H_{31}COOC_{30}H_{61} + H_2O \\ Palmitic\ acid & Myricyl & Myricyl\ palmitate \\ & alcohol & \end{array}$$

myricyl palmitate being the chief constituent of beeswax.

The classification and identification of fats is based upon the acids which they contain. Thus it is found that whereas beef suet and mutton fat consist chiefly of esters of the higher fatty acids, such as palmitic and stearic acids, butter contains a considerable quantity of the lower members of this same fatty series such, for example, as butyric, caproic, caprylic and capric acids; these acids, which are low boiling liquids readily volatile with steam, are known as volatile fatty acids and their presence or absence in a given sample of fat may be used for characterizing the fat. Thus, for example, the estimation of the amount of volatile fatty acid serves to distinguish genuine butter from its substitute margarine, which is relatively poor in volatile acids and contains chiefly higher fatty acids.

The more important members of the fatty acid series are given in the following list:—

НСООН	or CH ₂ O ₂	Formic acid*
CH₃COOH	$_{,,}$ $C_{2}H_{4}O_{2}$	Acetic acid
C ₂ H ₅ COOH	$_{1}$, $C_{3}H_{6}O_{2}$	Propionic acid *
C ₃ H ₇ COOH	$_{,,}$ $C_4H_8O_2$	Butyric acid

[&]quot;These acids do not occur in fats.

```
CH<sub>3</sub>>CH CH<sub>2</sub>CH<sub>2</sub>COOH or C<sub>6</sub>H<sub>12</sub>O<sub>2</sub>
                                                                    Isobutyl acetic or caproic acid
CH<sub>3</sub>(CH<sub>2</sub>)<sub>6</sub>COOH
                                             C_8H_{16}O_2
                                                                    Caprylic acid
CH<sub>3</sub>(CH<sub>2</sub>)<sub>8</sub>COOH
                                             "C10H20O2 Capric acid
                                             "C<sub>12</sub>H<sub>24</sub>O<sub>2</sub> Lauric acid
CH3(CH3)10COOH
CH<sub>3</sub>(CH<sub>2</sub>)<sub>19</sub>COOH
                                             ,, C<sub>14</sub>H<sub>28</sub>O<sub>2</sub>
                                                                    Myristic acid
                                             "C16H32O2 Palmitic acid
CH3(CH2)14COOH
CH<sub>3</sub>(CH<sub>2</sub>)<sub>16</sub>COOH
                                             ,, C_{18}H_{36}O_2 Stearic acid
,, C_{20}H_{40}O_2 Arachidic acid
CH2(CH2)28COOH
CH<sub>2</sub>(CH<sub>2</sub>)<sub>20</sub>COOH
                                             ,, CooH44Oo Behenic acid
```

It should be noted that these acids all conform to the general formula for the fatty acids, $C_n H_{2n} O_{2n}$ in which "n" may have any value, odd or even, but only those in which "n" is an even number are found to occur naturally in fats; the alleged occurrence in natural fats of acids with an uneven number of carbon atoms has in every case, so far recorded, been refuted on careful re-examination.

It appears probable, moreover, that all naturally occurring fatty acids have a straight and not a branched carbon chain, so that it is open to question whether the *iso*-butyl acetic acid which is said to have been found in fats was not, in reality, normal caproic acid of the formula $CH_3(CH_2)_4COOH$.

Besides acids of the fatty series whose general formula is $C_nH_{2n}O_g$, acids belonging to several other series, poorer in hydrogen than the above, are found in fats. The simplest example of such a series of acids is furnished by the acids of the Oleic series, the members of which differ from the corresponding members of the fatty acid series in having two atoms of hydrogen less.

Some of the more important acids of this group are given below.

I. Acids of the OLEIC or ACRYLIC series. General formula $C_nH_{2n-2}O_2$.

```
 \begin{array}{lll} C_5 H_8 O_2 & Tiglic \ acid \\ C_{18} H_{24} O_2 & Oleic \ acid \\ C_{18} H_{34} O_2 & Elaïdic \ acid \\ C_{18} H_{34} O_2 & Iso-oleic \ acid \\ C_{22} H_{42} O_2 & Erucic \ acid \\ C_{22} H_{42} O_2 & Brassidic \ acid \\ \end{array}
```

The most widely distributed of these acids is undoubtedly oleic acid, which, in the form of its glyceride triolein,

$$C_{17}H_{33}COOCH_{2}$$
 $C_{17}H_{33}COOCH$
 $C_{17}H_{33}COOCH_{2}$

forms an important constituent of most vegetable and animal oils.

- Acids of the LINOLIC series. General formula C_nH_{2n-4}O₂.
 - (a) Open chain compounds, $C_{18}H_{32}O_2$ Linolic acid and its various isomers. (b) Cyclic compounds, $C_{18}H_{32}O_2$ Hydnocarpic acid. $C_{18}H_{32}O_2$ Chaulmoogric acid.

3. Acids of the LINOLENIC series. General formula C_nH_{2n-6}O₂.

C18H20O2 Linolenic acid and its isomers.

4. Acids of the CLUPANODONIC series. General formula C_nH_{2n-s}O₂.

C18H28O2 Clupanodonic acid.

5. Acids of the RICINOLEIC series. General formula C_nH_{2n-2}O₃.

C18H24O3 Ricinoleic acid and its isomers.

The relationship between the five series of acids, which differ from each other successively by two atoms of hydrogen, as shown by the formulæ-

$$C_{n}H_{2n}O_{2},\ C_{n}H_{2n-2}O_{2},\ C_{n}H_{2n-4}O_{2},\ C_{n}H_{2n-6}O_{2},\ \text{and}\ C_{n}H_{2n-8}O_{2}$$

is similar to that subsisting between the three series of hydrocarbons having the general formulæ:--

$$C_nH_{2n+2}, C_nH_{2n}, C_nH_{2n-2}$$

The hydrocarbons of the first or Paraffin series are said to be saturated, by which is meant that each of the four valencies of their carbon atoms are fully satisfied, as shown by the following graphic formulæ :---

When, however, the graphic formulæ of the corresponding members of the second or Olefine series are written, it is found that if the tetravalency of carbon is maintained, there are not enough hydrogen atoms to satisfy all these valencies, and, in order not to leave any unsatisfied, the remaining valencies must be united to each other, thereby joining two carbon atoms to each other by more than one bond:—

In the next series of hydrocarbons, the acetylenes, by the loss of two more hydrogen atoms, the process has been carried a step farther, with the result that two carbon atoms are united by a triple bond:—

$$\begin{array}{ccc} HC \equiv CH & H - \stackrel{H}{C} - C \equiv CH \\ & \stackrel{|}{H} \\ Acetylene & C_{2}H_{2} & Allylene & C_{3}H_{4} \end{array}$$

All such substances containing two carbon atoms united together by more than one bond are said to be unsaturated, and are able to form additive compounds with many substances, notably the halogens.

Thus, while the saturated hydrocarbon will only react with chlorine or bromine by the replacement of one atom of hydrogen for each atom of halogen introduced into the molecule,

$$C_2H_6 + Br_2 = C_2H_5Br + HBr$$

Ethyl bromide

an unsaturated compound, such as ethylene, will add on the halogen directly—

$$C_2H_4 + Br_2 = C_2H_4Br_2$$

Ethylene
dibromide

the resulting additive compound being saturated.

It will thus be seen that it requires two atoms of bromine to saturate an unsaturated compound containing one double bond, and similarly it requires four atoms of halogen to saturate a compound containing two double bonds. In this way it is shown that since the acids of the oleic, linolic, and linolenic series require two, four, and six atoms of halogen respectively for saturation, they must contain respectively one, two, or three double bonds.

A measure of the degree of unsaturation of a given acid may accordingly be obtained by determining how much bromine it will absorb; as, however, the interaction with bromine is liable to be violent it is found more convenient to employ iodine, which, in addition to being less violent in its action than bromine, is also easier to handle.

A description of the method employed in determining what is known as the "iodine value" of fats is given below (p. 29).

EXTRACTION OF FATS.

The isolation of fats from admixture with other substances may be effected by extraction by means of fat solvents.

The principle of the extraction is to treat the dried mixture with a solvent which will dissolve only the fat and leave the other substances unchanged. The solvents most commonly used for this purpose are ether, light petroleum, carbon tetrachloride and carbon disulphide, the two latter being used chiefly on a commercial scale.

It must be borne in mind that besides extracting fats, ether will also dissolve out lecithin, cholesterol, essential oils and the somewhat indefinite group of substances known as lipoids.*

Moreover, other substances which are of themselves insoluble in ether may become soluble in the presence of fats.

Whatever solvent is employed must be tested before use to see that it leaves no residue on evaporation and is free from moisture.

A rough and ready method of extracting fat from a given sample is to place the finely divided and dried material on a filter paper folded into a funnel and to pour the fat-solvent on to it. The filtrate will contain most of the fat which may be recovered by evaporating off the solvent.

When it is desired to extract the fat quantitatively, the

^{*} This term has no chemical significance, and comprises a variety of different substances, which owing to their exhibiting the same solubilities as fats were called by Overton Lipoids (see p. 44).

operation is most conveniently carried out in a Soxhlet apparatus (see below).

Previous to extraction, the substance must be thoroughly dried. For this purpose it must either be gently heated in a current of dry air or else desiccated by means of alcohol or anhydrous salts.

The first method, which is the most convenient, should, however, be used with caution, as many fats may undergo chemical change during the process, as a result of which the material extracted by ether after drying may be very different from the substance originally present in the moist sample.

The second method, which consists in treating the sample to be dried with absolute alcohol for some hours and then filtering and pressing, depends on the fact that the alcohol withdraws the water without dissolving away any appreciable quantity of the fat; if treated two or three times in this way the substance will be practically free from moisture and can then be extracted under a Soxhlet with ether. The wet alcoholic filtrates on careful evaporation yield a residue which may be separately treated with ether to extract any fat contained in them. It is unnecessary to remark that the method cannot be employed if the fat to be extracted is soluble in alcohol

The third method of drying, which involves the use of anhydrous salts such as sodium sulphate, depends on the fact that the anhydrous salt when ground up with the moist tissue withdraws the water from it, forming the hydrated crystals. In a few hours the substance is sufficiently dry to be powdered. The chief objection to this process is the fact that a considerable bulk of salt has to be employed and consequently the volume of the material to be extracted is much increased.

PHYSICAL PROPERTIES OF FATS.

The naturally occurring fats vary in consistency from oils to wax-like solids; the solid fats have mostly a low melting point which is, however, rarely a sharp one, as natural fats are not simple substances, but are, as a rule, mixtures of several different chemical individuals; such mixtures never have sharp melting points.

All fats and fatty oils are lighter than water, their specific

gravity varying from about 0.900 to 0.970 at 15°. They are insoluble in water and at ordinary temperatures are sparingly soluble in cold alcohol, excepting castor oil which dissolves readily in this solvent; they, however, dissolve readily in ether, chloroform, petroleum ether, benzene, carbon tetrachloride or carbon disulphide,

CHEMICAL PROPERTIES OF FATS.

One of the most important chemical properties of fats is their decomposition by hydrolysis.

The term hydrolysis, which literally means loosening by water, is applied to any reaction in which a substance is broken up into two or more simpler ones with the fixation of water.

The following examples taken from a variety of different classes of compounds all illustrate this reaction:—

It will be seen from reaction (I) that the conversion of an ester into an acid and an alcohol is an example of hydrolysis, and since fats are esters it follows that they also must be capable of hydrolysis.

The reaction-

$$C_{17}H_{25}COOCH_2$$
 CH_2OH CH_2OH CH_2OH CH_2OH CH_2COOCH_2 CH_2OH CH_2COOCH_2 CH_2OH CH_2O

is, however, not readily brought about by water alone at ordinary temperatures; in the presence of enzymes, however, the hydrolysis may be effected at a moderate temperature with comparative ease (see p. 368).

The hydrolysis of fats for the purpose of preparing the *free* fatty acids may be effected in either of the following ways:—

1. By acting on the fat with superheated steam in the presence of a little lime or magnesia, which acts as a catalytic

agent.

This method is the one most commonly adopted by candle-makers for the preparation of fatty acids required in the manufacture of candles. The fat is subjected to the action of steam under pressure at 170° in large copper vessels in the presence of a small quantity of lime. The resulting mixture is then treated with sulphuric acid sufficient in amount to combine with the lime, after which the free fatty acids rise to the surface in a molten condition.

2. By the action of concentrated sulphuric acid.

The molten fats are stirred up in leaden vessels with 9 per cent of concentrated sulphuric acid, the mixture being heated to about 120° C. The mixture is then warmed with water to remove the acid, and the acids are further purified by distillation with steam.

SAPONIFICATION OF FATS.

Closely related to hydrolysis is the reaction known as saponification; this reaction, which literally means "soapmaking," is that which takes place when a fat is boiled with caustic alkali. The alkali acts in much the same way as water, breaking up the ester into glycerol and the fatty acid which, however, in this case, combines with the alkali to form a salt:—

$$\begin{array}{c|cccc} C_{17}H_{35}COOCH_2 & CH_2OH \\ \hline C_{17}H_{35}COOCH + 3HOH = 3C_{17}H_{35}COOH + \\ \hline C_{17}H_{35}COOCH_2 & Potassium stearate, \\ \hline C_{17}H_{35}COOCH_2 & CH_2OH \\ \hline C_{17}H_{35}COOCH_3 & CH_2OH \\ \hline \end{array}$$

It so happens that the sodium and potassium salts of palmitic, stearic, and oleic acids dissolve in water forming opalescent alkaline solutions which readily give a lather, and can, therefore, be used as soaps,* and hence the process by which

* The sodium and potassium salts of oleic acid and of the higher fatty acids, such as palmitic and stearic acids, when dissolved in water, are, to a large extent, hydrolysed into free fatty acid and caustic soda according to the equation:—

$$C_{17}H_{38}COONa + H_2O = C_{17}H_{38}COOH + NaOH$$

Sodium stearate Stearic acid

The stearic acid combines with some of the unhydrolysed soap to form an

they are made from fats is called saponification. Although alkali metal salts of other organic acids do not exhibit the characteristics of soap, the term saponification has nevertheless been extended to include all cases of the decomposition of an ester into the corresponding alcohol and the salt of the acid, even though that salt may have none of the characteristic properties of a soap.

The saponification of a fat on a small scale * in the laboratory may be effected as follows: boil the fat with about three or four times its weight of alcoholic potash under a reflux condenser. The alcoholic potash is prepared by dissolving caustic potash in about twice its weight of water and mixing the solution with twice its volume of alcohol. The heating should be continued until on pouring a little of the solution into a large volume of water an opalescent solution free from undecomposed fat results. The time required for this may vary from a few minutes to half an hour or more.

When the saponification is complete, the contents of the flask should be heated in an evaporating basin over a water

insoluble acid salt, giving rise to an opalescent or turbid solution. It is this insoluble acid salt which is responsible for the formation of a lather on shaking such a solution. The detergent or cleansing action of soap is dependent on the above reaction since the caustic soda detaches the greasy dirt which then becomes enveloped in a layer of soap solution from the lather and is so carried away.

In this connexion it is interesting to note the similar effect of soap on the formation of emulsions.

An emulsion may be defined as a mixture, under special conditions, of two otherwise immiscible liquids. Thus, for example, if olive oil is shaken up with water, the two liquids rapidly separate as soon as the shaking ceases. If, however, a little soap solution or some other substance such as gum acacia, tragacanth, saponin (see p. 183), or white of egg be added and the shaking repeated, an emulsion results owing to the oil particles being enveloped in a layer of soap or other substance which prevents their coalescing. Milk is an example of a naturally occurring emulsion; so also is latex contained in plants.

If pure olive oil, free from oleic or other acid, is shaken up with caustic soda no emulsion is produced; on the other hand, olive oil which has been kept some time and contains free oleic acid, when shaken up with caustic soda does produce an emulsion, thus showing that the emulsifying agent is not the free alkali but the soap produced in the second case from the soda and the oleic acid.

This may be also illustrated by Bütschli's experiment which consists in placing a drop of old olive oil containing 9 per cent of olcic acid on a little oro6 per cent aqueous solution of sodium carbonate. If examined under the microscope it will be seen to consist of a fine honeycomb structure, consisting of particles of oil, the whole apparently exhibiting amoeboid movements; these latter are due to difference in surface tension.

^{*} For commercial soap manufacture, see p. 5.

bath, and thoroughly stirred to get rid of the alcohol. If the free fatty acids are required sufficient sulphuric acid is then added to make the solution strongly acid, whereupon the fatty acids separate out and rise to the surface.

The aqueous layer contains the glycerol together with the excess of sulphuric acid and potassium sulphate.

CHOLESTEROL AND PHYTOSTEROL.

In addition to the trihydric alcohol glycerol, all fats contain a small quantity of the monohydric alcohols cholesterol and phytosterol* which form what is known as the *unsaponifiable* residue of fats.

These substances may be isolated from fats according to the following method devised by Kossel and Obermüller.†

An ethereal solution of the fat is mixed with a solution of sodium in alcohol; saponification takes place in the cold and the soap which is precipitated from solution can be filtered off; the filtrate, which is a mixture of alcohol and ether, contains the glycerol together with the so-called unsaponifiable residue consisting of phytosterol or cholesterol which may be obtained by evaporating the solvent.

The following method originally due to Allen and Thomson is recommended by Lewkowitsch for the estimation of the "Unsaponifiable Residue".

Five grams of the fat or oil are saponified by boiling under a reflux condenser with 25 c.c. of alcoholic potash containing II'2 per cent of caustic potash; when saponification is complete the alcohol is evaporated off and the residual soap is dissolved in 50 c.c. of hot water and transferred to a separating funnel of about 200 c.c. capacity, about 20-30 c.c. of water being used to rinse out the dish. After cooling, the mixture is shaken with 30-50 c.c. of ether and set aside until the ethereal layer has separated. (N.B.—The separation is accelerated by the addition of a little alcohol.) The soap solution is then run off from below into a second separating

^{*}The term phytosterol though employed by many authors to indicate a single definite substance is beginning to be used as a generic term for a whole group of closely allied substances the number of which is rapidly increasing as the investigation of vegetable fats proceeds.

[†] Kossel and Obermüller: "Zeit. physiol. Chem.," 1890, 14, 599; 1891, 15, 321.

funnel and shaken once more with a fresh quantity of ether. Two extractions should suffice, but it is safer to extract a third time. The ethereal extracts are then united, washed with a small quantity of water to remove any soap and transferred to a weighed flask; after evaporating off the ether, the flask is weighed again; the increase in weight gives the amount of unsaponifiable residue in 5 grams of the sample.

The isolation and identification of the unsaponifiable residue is of considerable importance for the purpose of establishing whether a given sample of fat or oil is of animal or vegetable origin, since animal fats all contain cholesterol while vegetable fats contain either phytosterol itself or a closely allied substance belonging to the group of phytosterols.

REACTIONS AND PROPERTIES OF CHOLESTEROL AND PHYTOSTEROL.

Cholesterol.

Cholesterol is a monohydric alcohol of the formula $C_{27}H_{45}OH$; its constitution is still unknown, although a great deal of work has been expended on this question. According to Windaus* it would appear to be a secondary alcohol containing an unsaturated group as expressed by the formula:—

Cholesterol occurs in the bile and forms the chief constituent of a certain type of gall stones; it also occurs in the brain and blood and is the chief alcohol constituent of wool fat.

It is insoluble in water and crystallizes from chloroform in needles and from ether or alcohol in rhombic plates, m.p. 148-150°. It may conveniently be obtained by extracting crushed gall stones with ether and evaporating the ethereal extract to dryness.

*Windaus: "Ber. deut. chem. Gesells.," 1912, 45, 2421; see also Dorée: "Biochem. Journ.," 1909, 4, 72.

Colour Reactions.—(I) Crystals of cholesterol pressed on a white porcelain surface and moistened with a drop of sulphuric acid (5 parts concentrated acid to I part of water) turn pink. The addition of a drop of dilute iodine causes a play of colours from red to blue or green.

(2) A solution of cholesterol in chloroform gently agitated with concentrated sulphuric acid turns red, while the sulphuric acid which forms the lower layer assumes a green fluorescence.

(3) On the addition of concentrated sulphuric acid drop by drop to a little cholesterol dissolved in a mixture of 2-3 drops of chloroform and about 10 drops of acetic anhydride, a transient pink colour is at first formed; on the addition of more acid, however, the colour changes to blue and finally to green.

Phytosterol or Sitosterol.

The term phytosterol was at one time employed to designate a definite chemical individual of the formula $C_{27}H_{45}OH$, but it is now used more as a generic term to include a number of different substances having certain properties in common. Thus Windaus and Hauth* showed that the substance obtained from Calabar beans and commonly known as phytosterol was in reality a mixture of two substances—(a) Sitosterol of the formula $C_{27}H_{45}OH$, and (b) Stigmasterol $C_{30}H_{47}OH$, an observation which has been confirmed by Salway.†

Similarly Klobb! describes a dextro-rotatory phytosterol of the formula C₃₁H₅₂O, 3H₂O occurring in *Anthemis nobilis* and a number of lævo-rotatory phytosterols of different formulæ obtained from *Matricaria Chamomilla*, *Tilia europaea*, *Linaria vulgaris*, and *Verbascum Thapsus*.§

All vegetable fats contain phytosterol, the amount varying from about 0.13 to 0.30 per cent and rising in the case of pea

^{*}Windaus and Hauth: "Ber. deut. chem. Gesells.," 1906, 39, 4378; 1907, 40, 3681.

[†] Salway: "Journ. Chem. Soc., Lond.," 1911, 99, 2154. ‡ Klobb: "Compt. rend.," 1911, 152, 327; "Ann. Chim. Phys.," 1911, viii.,

[§] See also Power and Rogerson: "Journ. Chem. Soc., Lond.," 1910, 97, 1951; Rogerson: "Amer. Journ. Pharm.," 1911, 83, 49; "Journ. Chem. Soc., Lond.," 1912, 101, 1040.

fat and the fat of Calabar beans to a considerably higher value.

Phytosterol crystallizes from alcohol in elongated plates and from ether in slender needles. The melting point of the pure substance varies somewhat according to the source from which it is prepared; it lies somewhere between 135 and 137° or it may be as high as 144°. The reason for this may be that the various substances obtained from different sources and described as one and the same substance are in reality different substances but all of a phytosterol nature. The colour reactions of phytosterol resemble those of cholesterol.

Cholesterol and phytosterol cannot with certainty be distinguished by means of their melting points, owing to the fact that phytosterol may melt at any temperature between 135 and 144° according to the source from which it is prepared. As, however, there is a considerable difference between the melting points of the acetates of these two substances the following procedure is recommended by Lewkowitsch.

The unsaponifiable residue remaining after evaporation of the ether (p. 15) is dried over a water bath and then dissolved in the least possible quantity of absolute alcohol and allowed to crystallize. The crystals which separate should be examined under a microscope; cholesterol crystallizes in four-sided plates and phytosterol in elongated hexagonal plates.

The alcohol is then evaporated off completely and the residue is carefully heated with 2 to 3 c.c. of acetic anhydride over a free flame until the liquid boils, the remaining acetic anhydride being evaporated off over a water bath. The residue is then re-crystallized two or three times from the least possible quantity of absolute alcohol, and the melting point of the crystals so obtained is determined.

Cholesterol acetate melts at 114.3-114.8°. Phytosterol acetate * melts at 125-137°.

SPONTANEOUS CHANGES IN FATS.

Rancidity.—Most fats when exposed to air and light sooner or later become rancid, acquiring an unpleasant taste and smell.

*The acetyl derivative obtained by Power and Moore from the root of Bryonia has the melting point 155-157°.

The actual cause of this change is as yet but little understood, though it appears probable that it is the result of the combined action of a number of different factors such as oxygen, light, moisture, bacteria and enzymes; the complex fats, and possibly also the small quantities of proteins and other impurities contained in them, are thereby broken down into simpler bodies such as the lower volatile fatty acids and aldehydes. It is frequently true that a considerable quantity of free acid is liberated in fats which have become rancid, and this is especially so in the case of fats such as butter which contain acids of low molecular weight, as butyric acid, the smell of which recalls that of rancid butter. It is, however, a fact that a fat may be acid without being rancid; cocoa butter, for instance, has usually an acid reaction but very rarely becomes rancid.

With regard to other constituents found in rancid fats, various authors have from time to time observed the presence of hydroxy-acids, aldehydes, alcohols, and of esters of lower fatty acids, but there appears to be a general consensus of opinion that glycerine does not occur.

Drying and Resinification.—Most fatty oils on exposure to the air tend to thicken, owing partly to polymerization and partly to oxidation; in some cases the oil actually dries up, leaving a more or less hard mass or a thin elastic film.

Those oils which only thicken, without actually becoming hard or dry, are called *non-drying* oils. They are composed for the most part of triolein (cf. p. 7) and contain only small quantities of solid fatty acids; to this class of oils belong the following: olive oil, almond oil, arachis or pea-nut oil, quince oil, cherry- plum- peach- and apricot-kernel oil, wheat-meal oil, rice, tea-seed oil, and hazel-nut oil.

Two further oils, namely castor oil and grape-seed oil, are also included in this group of non-drying oils, but they have a slightly different composition from the other members of this group. They are characterized by possessing a considerable percentage of glycerides of hydroxylated fatty acids, such as dihydroxystearic acid, a fact which is brought out clearly by their high acetyl values (p. 34).

In contrast with these non-drying oils are the so-called drying oils, among the more important of which are the follow-

ing: linseed oil, cedar-nut oil, hempseed, walnut, poppy-seed, and sunflower oil. These oils exhibit to a greater or less degree the tendency to absorb oxygen from the air, thereby drying up and leaving an elastic skin, a property which is made use of industrially in the manufacture of oil paints. These drying oils are composed chiefly of the glycerides of the unsaturated acids of linolic and linolenic series and contain only relatively small quantities of oleic acid. Owing to the large amount of unsaturated acids which they contain their iodine value (p. 29) is very high (120-200).

In addition to the above there is also a third group of vegetable oils known as the *semi-drying* oils whose iodine value and drying properties lie midway between those of the drying and non-drying oils. They differ from the true drying oils in containing no acids of the linolenic series and from the non-drying oils in containing linolic acid. The oils belonging to this category fall naturally into two sub-groups:—

- (1) The cotton-seed oil group, to which belong Soja-bean oil, maize oil, pumpkin, water-melon and melon-seed oils, beech-nut oil, cotton-seed, sesame and croton oils, and the lesser known oils of the apple, pear, orange, barley and rye seeds.
- (2) The rape oil group comprising garden cress, hedge mustard, wild radish, black mustard seed, white mustard seed, radish seed and rape or colza oil.

The oils of the latter sub-group have a lower saponification value (p. 29) than any other vegetable oils, and arachidic acid seems to be a normal constituent of them all.

To determine whether an oil is a drying one or not, a drop is spread on a glass plate, such as a microscope slip, and left for several days at atmospheric temperature. Non-drying oils such as olive and castor oils are unaltered after about eighteen days: semi-drying oils such as cotton-seed, sesame and rape oil are more or less dry, but still sticky in from seven to eight days, whereas real drying oils like poppy and especially linseed are quite dry in from three to six days.

GENERAL PROPERTIES AND REACTIONS OF FATS.

(I) All fats, both solid and liquid, are soluble in ether, light petroleum, carbon tetrachloride, chloroform and carbon di-

sulphide, but are only sparingly soluble in alcohol and insoluble in water.

(2) Being esters of glycerol they all contain this substance as may be proved by heating any fat with potassium hydrogen sulphate, whereby the glycerol is broken down into acrolein, which may be detected by its unpleasant odour.

$CH_2OHCHOHCH_2OH = CH_2: CHCHO + 2H_2O$

In order to show that the hydrolysis of fats gives rise to glycerol some fat should be saponified as described above and then acidified with hydrochloric acid; after filtering off the fatty acids, the filtrate is evaporated to small bulk over a water bath and the residue is extracted with alcohol, which dissolves out the glycerol leaving behind the salts. The presence of glycerol in the alcoholic extract may be proved by evaporating to dryness and applying the following test: two drops of the residue are carefully heated to about 120° with two drops of molten phenol and an equal quantity of concentrated sulphuric acid; the resulting resinous mass on cooling gives a brown solid which dissolves in ammonia forming a carmine-coloured solution.

- (3) All fats can be saponified by boiling with alcoholic potash. For this purpose 2 grams of fat may be boiled for fifteen minutes with 25 c.c. of 3 per cent alcoholic potash. The resulting mixture of potassium soap and glycerol is soluble in water; on acidifying the solution the free fatty acids are precipitated.
- (4) Fats leave a translucent mark on paper. Similar stains may be left by substances other than fats, but in most cases the stains disappear fairly rapidly as the substance evaporates.

SPECIAL TESTS FOR PARTICULAR CLASSES OF FATS.

Elaïdin Test.—This test, which is distinctive of the oleic series, depends on the fact that nitrous acid converts liquid olein into solid elaïdin, while it has no corresponding action on glycerides of linolic, linolenic, or isolinolenic acids. The test may be performed as follows:—

Ten grams of oil are shaken in a test tube with 5 grams of nitric acid (sp. gr. 1.38-1.41) and 1 gram of mercury for

three minutes or more until the mercury is completely dissolved. After the lapse of twenty minutes, the mixture is shaken for another minute, and is then set aside and the time noted which is required for the oil to solidify.

Olive oil requires one hour.

Arachis or earth-nut oil requires one and a half hours.

Colza and sesame oil require three hours.

Linseed oil gives a red pasty froth.

Hempseed oil remains unchanged.

The temperature of the mixture must be maintained constant during the test, and must not exceed 25°.

Bromide Test.—This test, which is chiefly used for distinguishing between drying and semi-drying oils, depends on the fact that linolic, linolenic acids and other unsaturated acids produce insoluble additive compounds with bromine, containing six or eight atoms of this element. According to Hehner and Mitchell * from 1-2 c.c. of oil are dissolved in 40 c.c. of ether containing a few cubic centimetres of glacial acetic acid; the mixture is then cooled to 5° and treated with bromine drop by drop until no more is absorbed. After three hours the precipitate is filtered off on a tared asbestos filter and washed four times with 10 c.c. of ether; it is then dried in a steam oven. The weight of the precipitate is directly proportional to the amount of unsaturated acids present in the fat.

Sulphuric Acid Test.—On mixing fats of the oleic series with concentrated sulphuric acid no heat is evolved, whilst with fats of the linolic series the opposite is the case.

COLOUR REACTIONS OF INDIVIDUAL FATS.

Many of the colour reactions described for fats are of doubtful value owing to the modifying influence of small quantities of resins and of proteins. The following tests are, however, fairly reliable:—

Badouin's Test for Sesame Oil.—Twenty c.c. of sesame oil are thoroughly shaken for a short time with 10 c.c. of hydrochloric acid (sp. gr. 1'9) containing 0'18 gram of cane sugar. A rose colour should appear immediately after the two layers of oil and water have separated; if left to stand longer the sugar solution causes a brown coloration.

^{*} Hehner and Mitchell: "The Analyst," 1898, 23, 313.

Solstien's Reaction for Sesame Oil.—Two or three volumes of oil or fat are dissolved in twice their volume of benzene (b.p. 70-80°) and gently shaken with three volumes of concentrated zinc chloride saturated with hydrochloric acid, the whole being kept immersed in a water bath at a temperature of 40° C.; when the zinc chloride has sunk to the bottom, the test tube is immersed up to the top level of the zinc chloride in a water bath at 80° C. If sesame oil is present a pink colour is produced.

Halphen's Reaction for Cotton-seed Oil.—Equal volumes of oil, amyl alcohol, and a one per cent solution of sulphur in carbon bisulphide are mixed in a test tube immersed to half its depth in boiling brine for about ten minutes. By the end of this time an orange colour should appear; if not add more carbon bisulphide and boil again. The colour is said to be due to the addition of sulphur to an unsaturated bond.

Sulphuric Acid Test.—Rape-seed oil and mustard oil when shaken with sulphuric acid (sp. gr. 1·53-1·62) produce grassgreen to blue-green colours. Linseed and hemp oil may also give similar colorations.

MICROCHEMICAL REACTIONS.

- 1. The microscopical appearance of oil when mixed with water is characteristic owing to its immiscibility with water and its different refractive index.
- 2. Its solubility in ether, chloroform, benzene, or other fat solvents is easily noted.
- 3. If oil be present in the preparation it will fairly rapidly turn brown and then black when treated with a one per cent solution of osmic acid. This is not absolutely conclusive since osmic acid stains proteins brown.
- 4. Tincture of alkannin, or a saturated solution of Scharlach R in 75 per cent alcohol, colours oil globules red or pink.

The reaction with the first-named reagent is often ill-defined and frequently fails when the alkanna used has been extracted from the root some time. The test is more satisfactory when freshly prepared tincture is used.

A similar reaction is given by Sudan III.

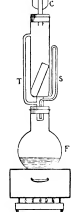
It is important to note that these and similar reactions are not conclusive of the chemical nature of the substances acted upon. For example, Sudan III not only stains oils red but also resins, latex, wax, and cuticle; chloroplasts are stained a pale red; cellulose, lignified walls, gelatinized membranes, starch, and tannin are unstained.

The staining tests mentioned above may be employed after extracting the oil with ether or other solvent.

QUANTITATIVE ESTIMATION OF FATS.

I. By Means of Soxhlet's Extraction Apparatus.—The fact that oils and fats are readily dissolved by ether, chloroform, and light petroleum, is made use of in their estimation; but it must be borne in mind that the method only yields correct results provided other substances, which would be extracted by the solvent employed, are absent from the material under examination.

The general arrangement of the apparatus required is given



in Fig. 1. The flask F, which is half-filled with the solvent to be employed, is connected to the extractor by a closely fitting cork. The material to be extracted is put into a thimble made of special quality filter paper and placed in the extractor, which is connected to a reflux condenser (C).

The method may be conveniently employed for determining roughly the proportion of oil in the reserve food of the castoroil seed, for example.

A number of seeds, freed from their testas, are carefully weighed and one by one are broken up in a perfectly clean glass basin with the well-rounded end of a glass rod. The material is then dropped into the thimble; any particles adhering to the basin or rod must be carefully removed by means of a platinum wire and also placed in the thimble.

The basin and glass rod should then be carefully washed with a few drops of ether to remove the last traces of fat, and the resulting solution should be added to the broken-up seeds in the thimble, care being taken not to employ enough ether for the solution to trickle through the thimble.

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The thimble is suspended in a steam oven for half an hour, and is then placed inside the extractor: a few small chips of porcelain are placed in the flask F, and the whole carefully weighed and then half-filled with freshly distilled ether. apparatus is then connected up. The ether in the flask F volatilizes and passes up the tube T into the extractor and condenser, and gradually fills the Soxhlet; on reaching a certain level it siphons over into the flask, carrying with it the fat in solution; once in the flask the ether is again vaporized and goes through the same process as before, the oil, however, remains behind. The ether is allowed to siphon off at least a dozen times,* and then, when most of the ether is in the extractor, the flask is disconnected. The ether in the flask is evaporated off and the flask is placed in a steam oven for half an hour, it is then allowed to cool in a desiccator and finally weighed.

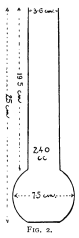
Per	cent	fat =	100	(y -	<u>z)</u>		
Weight of oil							y-z
Weight of flask and ch	ips						z
Weight of flask, chips	and o	il.					y
Weight of seeds .							X

If the ether has extracted substances other than fats, the result obtained will, of course, be too high. In such cases the ether extract may be saponified and the amount of fatty acid may be determined, from which the amount of fat originally present may be estimated.

2. By Saponification.—Apart from the fact that in some cases it is not possible to extract the fat quantitatively by Soxhlet's method with less than forty-eight hours' continuous extraction, the method is open to the objection that the substance must be dried previous to extraction, and this may involve loss or alteration of the fat; furthermore, the residue which is weighed as fat may not consist entirely of fat but may contain other substances which are extracted by the same solvents as the fats; this consequently necessitates a further examination of the residue for its saponification value, etc.

^{*} The number of times the liquid should be allowed to siphon off varies in every case. In order to ensure complete extraction the only safe method to adopt is to weigh the fat extracted after a certain time, then to attach the flask again and continue the extraction for some time longer and again weigh.

The following method which is due to Liebermann and Székely* has the advantage of giving in a short time a reliable value for the percentage of fat in almost any substance, and is specially convenient for the estimation of fat in fodder, meat, fæces, and physiological work in general. Five grams



of the sample are placed in a flask (of the dimensions given in Fig. 2) with 30 c.c. of 50 per cent caustic potash (sp. gr. 1.54). The mixture is boiled over a wire gauge for half an hour and frequently shaken. After cooling 30 c.c. of 90-94 per cent alcohol are added and the heating is continued for another ten minutes; the mixture is then cooled again and carefully mixed with 100 c.c. of 20 per cent sulphuric acid (sp. gr. 1.145) and thoroughly shaken after each addition; the temperature must be kept low so as to avoid any loss of volatile fatty acids. quite cold 50 c.c. of light petroleum (sp. gr. 0.6-0.7; b.p. about 60° C.) are added, and the flask is then closed with a tightly fitting rubber stopper and is thoroughly shaken for about ten seconds; the shaking is repeated about thirty times at intervals of one or two

minutes without removing the stopper. Saturated salt solution is then added until the lower aqueous layer reaches up to the 240 c.c. graduation which is marked on the neck. After shaking again a few times the flask is set aside in a vessel of cold water. When the petroleum containing the fatty acids in solution has separated, 20 c.c. are withdrawn by means of a pipette and are placed in a wide-mouthed 150 c.c. flask; 40 c.c. of 96 per cent alcohol, free from acid, are now added, together with 1 c.c. of a solution of phenolphthalein (made by dissolving 1 gram of accurately weighed phenolphthalein in 100 c.c. of 96 per cent alcohol) and the solution is titrated with N/10 alcoholic potash.

The titrated liquid is then carefully transferred in small portions at a time to a tared weighing bottle of about 80 c.c.

^{*} Liebermann and Székely: "Plüger's Archiv," 1898, 72, 360.

capacity, which is warmed over a gently boiling water bath; when the whole liquid has been evaporated to dryness, the residue is heated in an air oven for an hour at 100°, and, after cooling in a desiccator, is weighed with the glass stopper inserted to prevent the hygroscopic soap from absorbing any moisture from the air.

The amount of fat which corresponds to a given weight of soap may be calculated as follows:—

From the above equation it will be seen that in order to convert three molecules of soap into one molecule of fat three atoms of potassium, $3 \times 39^{\circ}I$, have to be withdrawn from three molecules of soap, and have to be replaced by 4I parts of $CH_2 \cdot CH \cdot CH_2$; this is equivalent to deducting $39^{\circ}I$ from one molecule of soap and adding $\frac{41}{3}$ or $13^{\circ}6$, or, in other words, deducting $25^{\circ}5$.

Hence, if "n" is the number of centimetres of N/10 caustic potash required for the titration, and since I c.c. N/10 KOH \equiv 00391 gram K \equiv 00136 gram C_3H_5 , we have to deduct from the weight of the soap W_8

$$n \times 00391$$
 and add $n \times 00136$

which is equivalent to deducting $n \times 0.0255$.

Also, since I c.c. of phenolphthalein solution on evaporation would leave o'o' gram of solid, this quantity must be deducted from the weight of the soap.

Hence the percentage of fat may be calculated from the relation

$$F = \left\{ \frac{W_\text{S} - \text{`or} - \text{(n \times `oo255)}}{\text{m}} \right\} \times \text{250}$$

in which "m" is the weight of the sample taken.

In estimating fat in flour or farinaceous grain by this method, it is best to subject the substance to a preliminary treatment by heating 5 grams of the sample for half an hour with 30 c.c. of dilute sulphuric acid (I:10), the mixture is then diluted with 50 c.c. of 50 per cent. caustic potash. Finally

the liquid is acidified with 60 c.c. of sulphuric acid (sp. gr. I·3) as described above. After the shaking with light petroleum is completed, 50 c.c. of 94 per cent alcohol are added instead of the salt solution; this has the effect of accelerating the separation of the petroleum layer which otherwise might take a long time.

Owing to the relatively small solubility of stearic acid in light petroleum the method may give too low a result in the case of substances very rich in stearin; the result should, therefore, be checked by a second estimation in which the number of shakings with petroleum are increased two or three fold. Leathes * has modified and considerably improved this method.

Kumagawa and Suto† have found that the following method gives good results: Two to five grams of the dry substance‡ are heated on a water bath for two hours with 25 c.c. of 5 N sodium hydroxide (20 grams in 100 c.c.) in a covered beaker. The mixture is then transferred to a separating funnel and acidified with 30 c.c. of 20 per cent hydrochloric acid. The fatty acids set free are taken up with ether, and the ethereal solution is filtered through asbestos and evaporated. The residue, which contains colouring matter, lactic acid and other substances as well as fatty acids, is dried for some hours at 50°, and then taken up with light petroleum, whereupon the impurities separate out in resinous form. After filtering through asbestos the petroleum is distilled off, and the residue, consisting of almost pure fatty acids, is dried at 50° to constant weight.

QUANTITATIVE METHODS EMPLOYED FOR THE CHARACTERIZATION OF FATS.

The following estimations are in common use for the commercial valuation of fats:—

(1) The Acid Number.

This is the number of milligrams of potassium hydroxide required for the neutralization of the *free* acids in a sample of fat.

* Leathes: "The Fats," London, 1910.

+ Kumagawa and Suto: "Biochem. Zeit.," 1908, 8, 212.

[‡]Yoshitaka Schimidzu ("Bioch. Zeit.," 1910, 28, 237) recommends using undried material since drying leads to a loss of fat, probably from oxidation.

This number is determined by dissolving 1 or 2 grams of the sample in 15 or 20 c.c. of a mixture of 1 part of alcohol with 2 parts of ether, and titrating the solution with N/10 alcoholic potash in the presence of phenolphthalein.

(2) The Saponification Value.

This is the number of milligrams of potassium hydroxide required for saponifying I gram of the fat.

From I to 2 grams of the sample are weighed out into a 250 c.c. conical flask; 25 c.c. of approximately seminormal alcoholic potash are then added, and the flask is attached to a reflux condenser and heated over a water bath for about half an hour; the solution is then diluted with 25 c.c. of water and cooled, then the excess of potash is titrated back by means of N/2 hydrochloric acid. In order to determine the strength of the alcoholic potash 25 c.c. of it are heated at the same time under exactly similar conditions in a second conical flask, but without any fat; in this way any error due to the effect of the alkali on the glass vessel is eliminated. The difference in the two titration readings gives the amount of acid equivalent to the potash used up in saponifying the fat, from which the number of milligrams of alkali required for I gram of fat may be calculated.

Since one molecule of any monobasic acid requires one molecule of potash, the magnitude of the saponification value is inversely proportional to the molecular weight of the acids contained in the fat.

			Molecular Weight.	Saponification Value.
Butyrin .			302	557:3
Palmitin.			806	208.8
Stearin .			8 9 0	189.1
Olein .			884	190'4
Coco-nut oil				246-260
Palm-kernel	oil *		_	242-250
Palm oil+				196-202
Olive oil.			_	185-196

(3) Iodine Value.

It was first observed by Hübl that an alcoholic solution of iodine containing mercuric chloride reacted at ordinary temperatures both with the free unsaturated acids and with

^{*} The oil contained in the kernel of the palm fruit.

[†] The oil contained in the fleshy part of the fruit.

their glycerol esters the fats. By elaborating the reaction, Hübl converted it into one of the most valuable criteria at present known for the detection and estimation of unsaturated acids in fats, and the so-called "iodine value" provides an excellent method of characterizing a fat.

For the determination of the iodine value of a fat the following solutions are required:—

- (a) An iodine solution made by mixing together equal volumes of two substances containing respectively 25 grams of iodine in water and 30 grams of mercuric chloride in 500 c.c. of 96 per cent alcohol. The two solutions should be mixed together about twenty-four hours before use, as the resulting mixture alters its strength considerably during the first few hours after it has been made.
- (b) A sodium thiosulphate solution containing roughly 24 grams of crystallized salt in 1 litre of water; the strength of this solution is accurately determined as follows: Twenty c.c. of a potassium bichromate solution containing 3.8657 grams of the pure salt dissolved in 1 litre of water are run into a stoppered bottle containing 10 c.c. of a 10 per cent solution of potassium iodide and 5 c.c. of concentrated hydrochloric acid. The resulting brown solution, if carefully made, should contain exactly 0.2 gram of iodine; it is at once titrated by means of the thiosulphate solution, and, supposing x c.c. were required to decolorize it then it follows that 1 c.c. of thiosulphate is equivalent to $\frac{O.2}{r}$ gram of iodine.
- (c) Chloroform or carbon tetrachloride, the purity of which should be tested by mixing 20 c.c. of it with 20 c.c. of the iodine solution and titrating the free iodine two or three hours after; the amount found should be exactly the same as that contained in 20 c.c. of the iodine solution to which no chloroform or carbon tetrachloride has been added.
- (d) A 10 per cent solution of potassium iodide made by dissolving 1 part of the iodide in 10 parts of water.
- (e) A starch solution freshly prepared by boiling up a suspension of 0.5 gram of starch in 50 c.c. of water.

The determination of the iodine value is carried out as follows:—

From 0.15 to 0.18 gram of a drying or marine animal oil,

0.2 to 0.3 gram of a semi-drying oil, 0.3 to 0.4 gram of a nondrying oil or 0.8 to 1.0 gram of a solid fat are accurately weighed from a weighing bottle by difference into a 500-800 c.c. bottle, provided with a well-ground stopper, and dissolved in 10 c.c. of the chloroform (c); 25 c.c. of the iodine solution (a) are then run in, and the stopper, which is moistened with potassium iodide solution (d) to prevent loss of iodine by volatilization, is inserted. If a clear solution is not obtained more chloroform must be added. The bottle is then left to stand in the dark and if the dark brown colour should disappear after two hours or less, another 25 c.c. of the iodine solution must be added, as it is essential that there should be a considerable excess of iodine. In the case of solid fats and nondrying oils the reaction can be considered as being complete after six to eight hours, but in the case of drying oils or fish oils twelve to eighteen hours are necessary. After the completion of this time from 15 to 20 c.c. of the potassium iodide solution (d) are added, and, after thorough shaking, the mixture is diluted with 400 c.c. of water. If a red precipitate of mercuric iodide is produced, more potassium iodide solution should be added. The excess of free iodine, part of which is dissolved in the chloroform and part in the potassium iodide solution, is then titrated by shaking with the standardized sodium thiosulphate solution until only a faint yellow colour remains. A little of the starch solution is now added and the titration is continued until the dark blue colour is destroyed. Twenty-five c.c. of the original iodine solution are then titrated in a similar way with the sodium thiosulphate, and the difference in the two results gives the amount of iodine absorbed. The amount of iodine thus absorbed by 100 grams of the fat gives the iodine value.

The values obtained by the Hübl method are generally considered to be very reliable and concordant, but the method is somewhat tedious, and for this reason the more rapid method of Wijs* is preferable.

The iodine solution required for this method is obtained by separately dissolving 9.4 grams of iodine chloride and 7.2 grams of finely powdered iodine in separate flasks in about 200

^{*}Wijs: "Zeit. anal. Chem.," 1898, 277; "Zeit. Unters. Nahr. Genussm.," 1902, 497.

c.c. of gently warmed glacial acetic acid. The two solutions are then united in a I litre graduated flask and made up to the mark with more glacial acetic acid.

This solution should be standardized on the following day by mixing 20 c.c. of it with 10 c.c. of 10 per cent potassium iodide solution and titrating the free iodine by means of the standard thiosulphate.

The actual determination of the iodine value is performed as follows:—

From 0.2-0.4 gram of fat should be carefully weighed and dissolved in 10 c.c. of pure carbon tetrachloride (which has been shown by a blank test not to absorb iodine); 25 c.c. of the iodine solution are then added and the flask is stoppered and set aside in the dark for one or two hours. The liquid is then transferred to a larger flask, the smaller flask being washed out thoroughly by means of 10 c.c. of potassium iodide solution and water until the total volume is about 300 c.c. The solution is then titrated with the thiosulphate. The difference between this reading and the amount required by 25 c.c. of the iodine solution is a measure of the iodine absorbed by the amount of fat.

The values obtained by Wijs's method are, as a rule, rather higher than those obtained by the Hübl method.

Appended is a list of iodine values of some important fats.

	• •						
(a)	DRYING OILS-						
	Linseed oil .						173-201
	Hemp-seed oil						148
	Sunflower oil						119-135
	Pine-seed oil	•	•	•			101-103
(b)	SEMI-DRYING OILS-						
	Beech-nut oil						104-111
	Cotton-seed oil						108-110
	Sesame .						103-108
	Rape oil (colza)		•	•			94-102
(c)	Non-Drying Oils-						
` '	Almond oil .						93-97
	Olive oil .						79-88
	Grape-seed oil						96-142
	Castor oil .				•		83 -9 0
(d)	VEGETABLE FATS-						
	Cacao butter						32-41
	Palm-kernel oil *						13-17
	Coco-nut oil *						8-10

^{*} Though described as oils these substances are both solid at ordinary temperatures, melting at about 25°.

(4) The Reichert Meissl Value.

This represents the number of cubic centimetres of N/10 caustic potash required for neutralizing the volatile acids liberated from 5 grams of a sample of fat under certain special conditions.

The determination is carried out as follows: Five grams of the sample are weighed into a 200 c.c. flask and saponified by warming with 70 c.c. of 10 per cent alcohol and 2 grams of caustic potash. The excess of alcohol is then evaporated off and the residue, after dissolving in 100 c.c. of water, is acidified with 40 c.c. of sulphuric acid (1:10); a few chips of asbestos are then dropped into the flask and the liquid is distilled through a Liebig condenser at such a rate that exactly 110 c.c. of distillate pass over in an hour. 100 c.c. of the distillate remaining after filtration are titrated with N/10 caustic potash in the presence of phenolphthalein. Appended are the numbers obtained for several different fats:—

Palm-oil	5-6.8	Lard .	0.68
Coco-nut oil	6.6-7.0	Tallow .	0.2
Linseed oil	0.0	Goose fat	0.5-0.3
Olive oil	o·6	Butter fat	20.6-33.1

The determination of the Reichert Meissl value is of considerable value for the detection of adulteration in butter, since any adulterant will at once lower the value.

(5) The Acetyl Value.

This is a measure of the amount of hydroxyl groups which a fat contains; its value depends upon the fact that compounds containing an alcoholic hydroxyl group react with acetyl chloride or acetic anhydride so as to replace the hydrogen of the hydroxyl by the acetyl group (CH_3CO-) as shown by the equation:—

$$ROH + CH_3CO > O = ROCOCH_3 + CH_3COOH$$

If the resulting acetyl derivative is saponified by means of caustic potash it breaks up as follows:—

$$ROCOCH_3 + KOH = ROH + CH_2COOK$$

and it is possible to determine the number of milligrams of

caustic potash which are thus utilized in combining with the acetyl groups to form potassium acetate,

The number of milligrams of potash required by the acetyl derivative obtained from I gram of the fat is termed the acetyl value of that fat.

Castor oil and grape-stone oil have particularly high acetyl values owing to the large proportion of hydroxyacids which they contain.

The following are the acetyl values of some of the more important oils, fats and waxes:—

Linseed oil .	3.98	Castor oil .	153-156
Olive oil .	10.64	Grape-seed oil	144
Rape-seed oil	14.7	Carnauba wax	55 24
Palm oil .	18.0	Lard	2.6
Palm-nut oil .	1.9-8.4	Butter	1.9-8.6

The following method, due to Lewkowitsch, has been adopted as the standard process.

About 10 grams of the fat are boiled in a round-bottomed flask under a reflux condenser for two hours with twice their weight of acetic anhydride. The mixture is then poured into a litre flask and boiled for half an hour with 500-600 c.c. of water, a slow stream of carbon dioxide being conducted into the liquid all the while to prevent bumping. After cooling, the upper layer of water is siphoned off and the lower oily layer is again boiled with water as above, the whole process being repeated three times. The oil is finally filtered and washed on the filter paper with boiling water until the filtrate is no longer acid, whereupon it is dried in an oven and weighed.

About 5 grams of the acetylated product are next saponified by boiling with alcoholic potash * as described under the determination of the saponification value. The alcohol is then evaporated off, and the resulting soap is dissolved in water.

Dilute sulphuric acid (1:10) is then added in excess and the solution is steam distilled until 600-700 c.c. of water have passed over. The distillate is titrated with N/10 caustic potash using phenolphthalein as indicator; the number of cubic centi-

^{*} Prepared by dissolving about 32 grams of 90 per cent stick potash in the least quantity of water and diluting to 1 litre with 96 per cent alcohol; the solution should be filtered after standing for twenty-four hours.

metres required for neutralization multiplied by 5.61 and divided by the weight of fat taken gives the acetyl value.

PHYSIOLOGICAL SIGNIFICANCE OF FATS.

The great function of fats in the economy of the plant is connected with nutrition. They form one of the most important food reserves of plants, and as such may occur in vegetative or in propagative organs.

With regard to their origin in plants very little is known; they first appear as very small vacuoles in the protoplasm which eventually run together forming large drops.

In some cases oil has been described as owing its origin to the activity of elaïoplasts, which are colourless bodies of various shapes usually grouped around the nucleus, and, like other plastids, of a protoplasmic nature. They are, or have been, supposed to act with regard to oil formation much as leucoplasts do with respect to starch formation. Elaïoplasts have been observed in many Monocotyledons such as Vanilla, Funkia, Gagea, Ornithogalum, etc., in the flower of a Dicotyledon, Gaillardia Lorenziana, and in Psilotum.

The development of the elaïoplasts of *Gaillardia* has been followed by Beer,* who found that they are formed by the aggregation of chloroplasts which then degenerate and give origin to the oil. He considers it is most unlikely that elaïoplasts perform any function of direct importance to the life of the plant, although they may in some cases, the corolla-hairs of *Gaillardia*, for instance, serve a biological purpose.

But although elaïoplasts may not perform the function originally ascribed to them, it does not necessarily follow that fats, more especially when occurring in the green parts of plants, may not be direct photosynthetic products. Thus Fleissig considers that in the case of Vaucheria, a plant which contains an abundance of fat, this substance is a direct photosynthetic product comparable to the starch and sugar in ordinary green leaves. On the other hand, it is, of course, possible that the fats in such cases may have been produced by secondary changes in the original product of photosynthesis.

In many cases there can be but little doubt that fats are

produced from carbohydrates; the work of Schmidt,* Le Clerc du Sablon,† and others has shown that as the carbohydrates disappear so fats appear. For example, in the case of the almond the seeds when they begin to ripen are rich in carbohydrates and poor in fats, whereas the reverse is true when they are fully matured. The same holds true for the seeds of *Ricinus* and *Pæonia*. The nature of the carbohydrates used up in this process varies in different plants; thus it is stated that in the case of the olive mannite is the carbohydrate. This statement, due to de Luca, is not accepted by other investigators of the same plant; according to Funaro the mannite does not appear until after the oil has been formed.

In the case of *Ricinus* seeds the oil is formed from glucose, and in *Pæonia* principally from starch. The facts that fat may be translocated as such, provided it be an emulsion sufficiently fine, or in the form of fatty acid or glycerine, suggest that the fats in seeds have not been formed *in situ*, but have been conveyed there. This may be true to a certain extent, but consideration of the fact that fat will appear as the carbohydrates disappear in immature seeds removed from the parent plant, together with the facts relating to the formation of fats in vegetative organs under the influence of cold (p. 3), leads to the conclusion that the substances in question are formed at the expense of carbohydrates. Further, corroborative evidence is afforded by well-ascertained facts relating to similar problems in animals.

Ivanow, ‡ experimenting with rape seed, has shown that they contain a lipase which may either hydrolyse a fat or may synthesize one from fatty acid and glycerine. Thus, if a glycerine extract of the seed be mixed with oleic acid, fat is synthesized, but, on diluting with water, the fat is split up again. This same author § has published important observations on the synthesis of fats in oily seeds mainly from the carbohydrates glucose, sucrose, and starch. These substances are synthesized in the order given, the last two being first

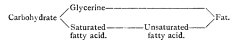
^{*} Schmidt: "Flora," 1891, 74, 300.

[†] Le Clerc du Sabion: "Compt. rend.," 1893, 117, 524; 1894, 119, 610; 1896, 123, 1048; "Rev. Gen. Bot.," 1895, 7, 145; 1897, 9, 313.

[‡] Ivanow: "Ber. deut. bot. Gesells.," 1911, 29, 595.

[§] Ibid., "Beith. bot. Centr.," 1912, 28, 159.

hydrolysed. The initial acids to be formed are characterized by a low iodine value, showing that they are saturated. Further, since the Reichert Meissl value is constant and does not vary with the acid number, it is concluded that the acids first formed belong to the higher members of the The saturated acids are followed by the unfatty series. saturated. Ivanow gives the following scheme to indicate the essential stages in the synthesis of fat in a typical instance such as the seed of flax :---



During the germination of oily seeds a reversal of this process takes place. The work of Schmidt, Green,* Le Clerc du Sablon, and others, has shown that the first process is that of hydrolysis which splits the fat into a fatty acid and glycerine, lipase being the active agent.

Thus in the sunflower Miller + found that less than I per cent of free fatty acid was present in the oil of the cotyledons of the resting seed; as germination proceeded there was a gradual increase, thus the ether extract of the cotyledons of a seedling in which the plumule was just showing contained 30 per cent of fatty acid.

The presence of the acid may be demonstrated in such germinating seeds, but the same statement does not hold for glycerine, probably because it is translocated with great rapidity, and is quickly transformed. There can, however, be no doubt that this substance is formed because if, for example, castor oil be subjected in vitro to the action of lipase obtained from Ricinus seeds, the presence of glycerine may be detected with ease.

With regard to other changes which the original fat undergoes during germination, Schmidt found that the iodine number of the unsaturated acids and oils decreased during germination, which indicates that saturation of the acid radicles takes place. This is controverted by von Fürth, ‡ who found

^{*} Green: "Proc. Roy. Soc., Lond.," 1890, 48, 370.

[†] Miller: "Ann. Bot.," 1910, 24, 693. ‡ Von Fürth: "Hofm. Beitr. Chem. Phys. Path.," 1904, 4.

no change in the iodine value. The observations of Schmidt, however, have been corroborated by Miller, who found that in *Helianthus annuus* the iodine value decreased from 136.2 for the seed to 67.4 for a seedling with the plumule just elongating.

Further corroboration is given by Ivanow* who, for his study on the transformation of fats during germination, selected flax, hemp, rape, and poppy seeds, since each is characterized by the possession of fats rich in acids of a specific series. Thus the oil of hemp seed is rich in acids of the unsaturated linolenic series, whilst poppy-seed oil is rich in acids of the saturated fatty acid series.

By ascertaining the iodine and other values of the fats of these seeds at different periods of germination, it was found that the acids disappeared in the sequence linolenic, linolic, oleic, and, finally, palmitic; in other words, the acids were consumed at a rate inversely proportional to their degree of saturation.

Ivanow considers that the fall in the iodine value of the fats is due rather to the rapidity with which the more unsaturated fatty acids are used up in the formation of carbohydrates rather than to their oxidation. He further found that the saturated fatty acids not uncommonly exist in a free state whilst the unsaturated acids occur in the form of glycerides.

Von Fürth also found that during germination the acetyl value decreased from 87.5 in the resting seed to 50.5 in the young seedling, from which he concluded that the normal fatty acid does not change into hydroxy fatty acid. Also, he could find no proof of the fatty acid breaking down into simpler substances as indicated by the molecular weight remaining practically constant.

This hydrolysis is the first action, but it is not the final one since carbohydrates quickly appear during the germination of such seeds. Since the days of de Saussure, who was the first to draw attention to this phenomenon, much evidence relative to this carbohydrate formation has accumulated.

In the case of *Ricinus* le Clerc du Sablon found that the resting seed contained 69 per cent of oil and 4 per cent of sugar, but in a seedling 11 cm. high the oil had fallen to 11 per

cent and the sugar had risen to 14 per cent. It was further found that the sugar contained in the resting seed has a slight excess of non-reducing sugar, which increased more rapidly than the reducing sugar; finally, however, the latter variety preponderated.

Le Clerc du Sablon also found the same relation between oil and sugar to obtain during germination of rape, hemp, poppy, almond, and walnut.

Similar observations have been made by Green and Jackson,* who found that in the resting seed of *Ricinus* the most abundant sugar is sucrose, which gives place to invert sugar in the early stages of germination. Subsequently the sucrose increases in amount, and occurs in quantities greater than the invert sugar; thus there is reason for supposing that the sucrose is a temporary reserve food.

The following table which summarizes the changes in the sugar content is taken from Green and Jackson's paper:—

Time of germina-	Invert sugar	Cane sugar
tion in hours.	in milligrams.	in milligrams.
0	1·1	10.7
45	2·7	5.17
69	2·3	0
117	6·7	19.4
168	5·2	10.5
216	19.2	35.8
240	29.01	35.8
312	40.8	52.6

Miller has found that in the sunflower, Helianthus annuus, the amount of ether extract of the cotyledons diminishes gradually from the beginning of germination, the most rapid depletion occurring during the period between the first appearance of the seed-leaves above ground and the point of full expansion. Also, the greatest increase in the hypocotyl and roots coincides with the period of maximum depletion from the seed-leaves. With regard to the sugar content, Miller states that the resting embryo contains about 4 per cent of sucrose, during germination there is a decrease, and this is followed by a gradual increase until the seed-leaves begin to unfold. Up to this stage the cotyledons contain only a non-reducing sugar, but as

^{*} Green and Jackson: "Proc. Roy. Soc., Lond.," B., 1906, 77, 69.

the seed-leaves assume the functions of foliage leaves a reducing sugar appears, and, in a short time, is the only sugar present. In the hypocotyl and roots the amount of sugar rapidly increases until in seedlings about 4 inches long it may amount to 20 per cent of the dry weight, then a gradual decrease takes place. There is also a small increase in the amount of starch

The nature of the carbohydrate differs in different plants; thus in addition to the above-mentioned plants, during the germination of *Allium* and of *Cucumis* much glucose makes its appearance; this is also true, although to a lesser degree, for *Cannabis sativa*, in which case the glucose is quickly transformed into starch.

In other instances starch is said to be the carbohydrate formed.

It is thus seen that there is an intimate connexion between carbohydrates and oil, and the question naturally arises how is the one connected into the other, to which there is no answer.

The consideration of the formulæ of the substances in question shows that fats poor in oxygen give rise to carbohydrates rich in oxygen, and vice versa; but as to how this is accomplished nothing of a definite nature is known.

Many suggestions have been put forward, and before mentioning these the reader may be reminded of the large amount of oxygen which is absorbed during the germination of oil-containing seeds.

Detmer considered that starch may arise from the free oleic acid according to the equation:—

$$C_{18}H_{34}O_3 + 27O = 2 (C_6H_{10}O_5) + 6CO_2 + 7H_2O$$

According to Maquenne the sugar has an origin depending upon the nature of the oil; thus if the fat be saturated, of its hydrolytic products the glycerine gives rise to the sugar, whilst the fatty acids are used up in oxidative processes. If, on the other hand, the fat be unsaturated, the fatty acid contributes to the formation of the sugar.

This change is supposed to be effected by the oxidation of the chain at the double bond setting free two unsaturated groups which by polymerization give rise to sugar.

These conclusions are based on the observations that during the germination of the seeds of *Arachis* the carbohydrate increases to 5.6 per cent of the dry weight, whilst in *Ricinus* the increase is 10 per cent. The glycerine of the fat would be sufficient to form about 5 per cent of carbohydrate; this roughly was the amount observed in the case of *Arachis*, but in *Ricinus* the amount was about three times as great.

Mazé has put forward the suggestion that the transformation of oil into sugar is effected by an enzyme.

It has already been mentioned that glycerine so far has not been demonstrated in germinating fatty seeds; this may be owing to its powers of rapid diffusion or to the fact that it is used up in the synthesis of other substances. it has just been mentioned that Maquenne thought that it might be the origin of the sugar in some cases at any rate; Green originally thought that such was the case in Ricinus, an opinion which he no longer holds. Green and Jackson state that there is reason to suppose that the protoplasm of the endosperm of *Ricinus* is increased at the expense of the initial reserve food-materials; subsequently, further carbohydrates for the nutrition of the embryo are formed by the activity of this protoplasm: in other words, these authors do not consider that the increase in sugar during germination and the decrease in oil are directly associated: the disappearance of the latter and the formation of the former are "features of a new metabolism set up in the cells as germination becomes established". Also they express the opinion that the glycerine may be used up in the formation of lecithin.

Le Clerc du Sablon has put forward the idea that there might be present an enzyme which acts on the fat without liberating the glycerine.

These views are concerned chiefly with the formation of carbohydrates from fats; a reversal of the process might or might not explain the formation of fats from carbohydrates.

The whole question is of considerable difficulty and, of course, refuge may be taken in the hypothesis first put forward by Nägeli that the fats are products of the disintegration of the protoplasm. Thus the carbohydrates might be assimilated by the protoplasm which might produce the oil by some catabolic process.

With regard to the possible formation of fats from proteins very little information is available. On the animal side there is some evidence to show that substances derived from proteins may be so utilized; a possible connexion may be found in the phospholipines (phosphatides) which are compounds of fatty acids containing either nitrogen or phosphorus, or both.

Leathes* points out that the fatty acid may be formed from glucose by processes analogous to the synthesis of butyric acid from lactic acid which in turn is formed from the glucose. For the underlying reasons, which are rather too complicated to be dealt with here, Leathes's monograph must be consulted. It may, however, be pointed out in this connexion that the investigations of Hanriot are very significant; he found that, in attempting the oxidation of fat *in vitro*, 15 per cent of its weight of oxygen was absorbed, and in the products of its oxidation butyric and acetic acids occurred, but no carbohydrate.

In conclusion brief mention may be made of Schmidt's views regarding the translocation of fats. He considers that in many cases the oils may be transported as such to those organs requiring it, for he found that the amount of fatty acid present in the germinating seeds was smaller than would be supposed if it were hydrolysed before translocation, also that neutral oil appears in regions of the plant removed from the storage organ.

He considers the walls of cells are permeable to oil; provided it be an emulsion sufficiently fine, and especially if a free fatty acid be present, the permeability being directly proportional to the amount of such acid present. It is thought that the acid forms a soap in the walls, and thus facilitates the passage.

It is not improbable that both methods are adopted by the plant, viz. the translocation of the products of the dissociation of the fat, and the translocation of oil *qua* oil.

^{*} Leathes: "The Fats," London, 1910.

WAXES.

The chief function of waxes in plants is to form a protective covering against undue evaporation of water. They are found most commonly in or on the cuticle of leaves and fruits where they give rise to the glaucous effect.

As already stated, the waxes resemble the fats in their chemical constitution in so far as they are esters, but they differ in the nature of their alcohol constituent which is not glycerol but is usually a monohydric alcohol such as cetyl alcohol $C_{16}H_{33}OH$, carnaubyl alcohol $C_{24}H_{49}OH$, pisangceryl alcohol $C_{24}H_{49}OH$, ceryl alcohol $C_{26}H_{53}OH$, myricyl alcohol $C_{30}H_{61}OH$, cholesterol or phytosterol $C_{27}H_{45}OH$.

In addition to the acids already mentioned as occurring in fats, the following are also met with in waxes in the form of esters: ficocerylic acid $C_{13}H_{26}O_{2}$, carnaubic acid $C_{24}H_{48}O_{2}$, and pisangcerylic acid $C_{24}H_{48}O_{2}$, as well as acids belonging to series of the general formula $C_{n}H_{9n-2}O_{2}$ and $C_{n}H_{9n}O_{3}$.

The term wax used in the chemical sense has reference only to the chemical composition of these substances, regardless of their physical state of aggregation, and consequently both liquid and solid waxes are known.

Waxes of the former class are, however, only known in the animal kingdom, they are ordinary sperm oil and arctic sperm oil.

Among the better-known vegetable waxes may be mentioned:—

- (a) Carnauba Wax obtained from Copernicia cerifera; this wax contains ceryl and myricyl alcohols, and two acids, cerotic acid $C_{26}H_{52}O_2$, and carnaubic acid $C_{24}H_{48}O_2$, together with a hydroxy-acid of the formula $C_{21}H_{42}O_3$.
- (b) Pisang Wax obtained from the leaves of Cera musae is the pisangceryl ester of pisangcerylic acid.

The following are some of the more important waxes of animal origin:—

Wool wax, better known as wool fat or lanolin (which is rich in cholesterol), beeswax, spermaceti, and Chinese insect wax.

PHYSICAL AND CHEMICAL PROPERTIES OF WAXES,

Waxes are soluble in all the ordinary fat solvents such as benzene, ether, chloroform, etc., though they are rather less soluble than the fats.

Being free from glycerides the waxes, when heated, give no smell of acrolein; they do not become rancid like the fats, and are less easily hydrolysed, but they can be decomposed by prolonged heating with alcoholic potash.

FURTHER REFERENCE.

Lewkowitsch: "Chemical Technology and Analysis of Oils, Fats, and Waxes," London, 1915.

PHOSPHATIDES, LECITHINS, OR PHOSPHOLIPINS.

Closely related to the fats is the group of substances known as phosphatides, phospholipins or lecithins, the last name being derived from the Greek $\lambda \epsilon \kappa \iota \theta_{05}$, meaning egg yolk, from which substance the first representative of the class was prepared.

The name phosphatide was first given by Thudichum to a number of substances containing phosphorus which he obtained from brain. Subsequently Overton introduced the term lipoid to represent a group of substances occurring in the animal tissues which resembled fats in their solubilities. Leathes uses the term phospholipins, in place of phosphatides, to denote compounds of fatty acids containing phosphorus and nitrogen, and proposes the name of lipins for compounds of fatty acids that contain nitrogen but no phosphorus.

OCCURRENCE.

Lecithin compounds occur in the grains of cereals, in the seeds of several Leguminosæ, *Ricinus*, and species of *Pinus*; in the leaves of *Castanea*, and in Fungi; they are also widely distributed in animals. In fact, these substances are stated to occur in small quantities in all living cells, and they appear to be more especially abundant where fats occur.

The approximate amount of lecithin contained in various substances may be seen from the following table:—

Egg yolk					9.4 per cent
Liver .					2'I ,,
Blood .					1.8 ,,
Leguminous	see	ds			o·8-1·64 per cent
Cereals .					0*25-0*53 ,,

Botanically, the term lecithin is generic, and plant products so called have not yet been obtained in a pure state and contain other substances of a similar nature.

PREPARATION.

The most convenient source for the preparation of lecithin is egg yolk. This substance is extracted with five times its volume of 96 per cent alcohol; the extract is then cooled to o°, filtered and precipitated with an alcoholic solution of cadmium chloride; the precipitated double salt is next washed with alcohol and ether; it is then decomposed by boiling with eight times its quantity of 80 per cent alcohol and carefully adding a concentrated solution of ammonium carbonate until all the cadmium is thrown out of solution; the solution is filtered whilst hot and on cooling the filtrate to 10° the lecithin is deposited. It may be purified by dissolving in chloroform and precipitating from solution by the addition of acetone in which lecithin is insoluble.

Pure lecithin has not as yet been obtained from vegetable sources, the substances isolated by Winterstein* and his collaborators from wheat flour and from the seeds of Avena sativa, Lupinus albus, L. luteus, Vicia sativa, from the leaves of Æsculus hippocastanum, etc., being mixtures which, moreover, contain a carbohydrate complex. For an account of the methods employed in the extraction of these substances the original papers should be consulted. Smolensky † found that wheat germs (i.e. the embryos which are a bye-product of the flour mills) yielded a phosphatide whose composition was much closer to that of ordinary lecithin than was that obtained from the flour.

^{*}Winterstein and Hiestand: "Zeit. physiol. Chem.," 1907, 54, 288; Winterstein and Smolensky: id., 1908, 58, 506; Winterstein and Stegmann: id., 1908, 58, 527. See also Schulze and Likiernik: id., 1891, 15, 405; Schulze: id., 1895, 20, 228.

⁺ Smolensky: id., 1908, 58, 522.

REACTIONS AND CHARACTERISTICS.

The following are some of the more characteristic reactions:—

- If to an alcoholic solution of lecithin an alcoholic solution of cadmium chloride be added, a white precipitate of the cadmium chloride double salt is formed.
- 2. If a little lecithin is boiled with caustic soda, trimethylamine is formed, and may be identified by its characteristic smell; the solution contains sodium salts of fatty acids; on acidifying with sulphuric acid the fatty acids are precipitated.
- 3. Lecithin gives a purple coloration when added to a mixture of strong sulphuric acid and sugar solution.

The lecithins are yellow or yellowish-white wax-like solids with a peculiar odour; they are very hygroscopic, but some of them when carefully dried in a vacuum can be obtained in form of powder. They dissolve readily in the ordinary fat solvents, such as ether, chloroform, carbon tetrachloride, benzene, carbon disulphide, and also in oils and fats; they are also soluble in hot alcohol and ethyl acetate, but are only sparingly soluble in acetone and methyl acetate, so that these two substances may be used for the purification of the crude product. They are precipitated from alcoholic solutions by alcoholic solutions of platinic or cadmium chlorides.

When mixed with a small quantity of water they swell up, forming slimy threads, known as myelin forms; with excess of water they "dissolve," forming colloidal solutions which are not coagulated by boiling, but from which they may be precipitated by the addition of certain salts, such as those of barium and calcium.

As already stated, phosphatides dissolve in the same organic solvents as the fats, and are consequently liable to be extracted from the tissues together with fats; this fact must be borne in mind in estimating the amount of fat in any substance by the method of weighing the residue remaining after the evaporation of an ether extract.

The chemical composition of the phosphatides * differs from that of the fats primarily in containing the two elements nitrogen and phosphorus in addition to carbon, hydrogen,

^{*} See Maclean: "Biochem. Journ.," 1915, 9, 351.

and oxygen. According as they contain one or two atoms of phosphorus in their molecules, they are classed as monoor di-phosphatides. The lecithins, like the fats, are esters of glycerol with higher fatty acids, such as palmitic, stearic, and oleic acids, but they differ from the fats in being at the same time esters of phosphoric acid, as is shown by the following formula of lecithin from egg yolk:—

Lecithin is readily hydrolysed by boiling with alkalis, notably baryta, and is also broken up by lipase, and, less readily, by mineral acids. The products of its hydrolysis are glycero-phosphoric acid:

$$CH_2OHCHOHCH_2OP = (OH)_2$$

choline HON(CH₃)₃CH₂CH₂OH and fatty acids; a similar hydrolysis takes place in the germinating seed.*

Originally it was considered that the fatty acids of lecithin were either stearic, palmitic, or oleic, but it has been found that the iodine values of the acids obtained from lecithin are much higher than would be given by these acids alone.

CHOLINE.

To examine the products of the hydrolysis of lecithin, this substance is heated with a solution of barium hydrate in excess; a baryta soap is formed, which may be filtered off. The aqueous solution contains barium glycero-phosphate and choline; the latter may be extracted as follows.†

Treat the solution with a stream of carbon dioxide until no more barium carbonate comes down. Filter and evaporate the filtrate to dryness. Treat the residue with absolute alcohol, which will dissolve the choline but not the barium glycerophosphate. The alcoholic solution, if treated with an alcoholic

^{*} Schulze: "Z. physiol. Chem.," 1887, 11, 365; Schulze and Frankfurt: "Ber. deut. chem. Gesells.," 1893, 26, 2151.
†Leathes: "The Fats," "Monographs of Biochemistry," London, 1910.

solution of platinic chloride, gives a precipitate of the double platinichloride of choline.

Green and Jackson* give the following method: Allow the finely divided material to stand for some days under absolute alcohol. Pour off the extract, and evaporate to dryness; the residue is again extracted with absolute alcohol, and finally with a mixture of alcohol and ether. These extracts are mixed, and the solvents evaporated off. The choline is contained in the residue. The following tests may be employed for its detection:—

I. Boil a strong aqueous solution; decomposition ensues and trimethylamine is given off, which may be recognized by its fish-like smell.

$\begin{array}{ll} HON(CH_3)_3CH_2CH_2OH = OHCH_2CH_2OH + N(CH_3)_3 \\ Choline & Trimethylamine \end{array}$

- 2. Add platinic chloride to the aqueous solution; a double platinum salt is formed, which crystallizes on standing. The crystals are soluble in 15 per cent alcohol. Should the crystals not appear, proceed as follows:—
- 3. Dissolve choline in alcohol and add an alcoholic solution of platinic chloride. Filter off the yellow precipitate, wash with alcohol and dissolve in as little water as possible. Place the solution in a watch glass, and stand in a desiccator. Hexagonal plates will be deposited.
- 4. In order to detect very small quantities, Rosenheim recommends the following method.† Prepare the double platinum salt, place a drop or two on a glass slip, and allow to evaporate. Add a drop of a solution containing 2 grams of iodine and 6 grams of potassium iodide in 100 c.c. of water, and examine under the microscope. Dark brown prisms or plates will appear and then disappear as evaporation takes place; they will reappear on adding another drop of iodine solution.

Choline is both a tertiary amine and an alcohol; the aldehyde and acid corresponding to it are both known.

 $\begin{array}{ccc} HON(CH_3)_3CH_2CH_2OH & HON(CH_3)_3CH_2CH(OH)_2 & HON(CH_3)_3CH_2COOH \\ Choline & Muscarine & Betaine \end{array}$

^{*} Green and Jackson: "Proc. Roy. Soc., Lond.," B., 1906, 77, 69.

⁺ Rosenheim : " J. Physiol.," 1905, 33, 220.

The aldehyde, which goes by the name of muscarine, occurs in *Agaricus muscarius*; it is a powerful poison (see p. 275).

By the bacterial decomposition of choline another very poisonous base, neurine, may be obtained; this substance differs from choline by the elements of water.

$$HON(CH_3)_3CH_2CH_2OH$$
 $HON(CH_3)_3CH=CH_2$
Choline Neurine

The choline complex of lecithin on further decomposition can give rise to nitrogen bases, such as dimethylamine $HN(CH_3)^2$ and trimethylamine $N(CH_3)_3$; these substances, which have a fishy smell, also occur in herring brine and are probably there produced from a similar source. They have similarly been obtained from the leaves of *Chenopodium vulvaria*, from the blossoms of *Crataegus Oxyacantha*, from species of *Pyrus*, and from the seeds of *Fagus* (see p. 276).

Lecithins form compounds with sugar, and it is stated that all lecithins of a vegetable origin are in combination with carbohydrates; galactose, glucose and pectose having been identified. The amount of this combined sugar varies pretty considerably; it may be as high as 16 per cent according to Winterstein and Hiestand.*

Formation of Lecithin.

The following table, due to Green and Jackson,† shows the relation between the lecithin, fatty acid, and oil of the endosperm of *Ricinus*, expressed in per cent of weight of the seeds at different stages in their germination:—

Degree of development.	Oil in seeds.	Fatty acid in seeds.	Lecithin.
Resting seeds .	82.8	2°2	·236
Testa just cracked .	67.5	4°6	·17
Radicle protruding 1-2 cm.	52.5	11°9	·475
Root system established .	23.6	16°89	·873

From this it appears that lecithin is formed during germination; although there is, during the early stages of germination, a

^{*} Winterstein and Hiestand : loc. cit.

⁺ Green and Jackson: loc. cit.

diminution in the quantity present. It was found when once the maximum was reached that this amount remained constant until the whole of the endosperm was used up.

The products of the decomposition of lecithin, viz. a fatty acid, glycerophosphoric acid, and choline, have been detected by Green and Jackson in the endosperm of germinating seeds of Ricinus, and they suppose that the action is reversible, so that the legithin is formed by the combination of the products of decomposition of the oil and protein reserves of the seeds. Thus the oil provides the fatty acid and the glycerol, of which the latter combines with phosphorus, obtained from the aleurone grains, to form glycerophosphoric acid. The choline is provided by the decomposition of the proteins by means of a tryptic enzyme. But however this may be, in view of our ignorance of these substances, and the fact that vegetable lecithins apparently have seldom or never been obtained in a state of purity, and the uncertainty relating to some of their cleavage products, it does not appear profitable further to consider here the theories which have been advanced to explain their formation.

Physiological Significance.

Nothing of a very definite nature is known of the physiological significance of the lipoids. Overton points out that under certain conditions lecithin and similar substances have the power of absorbing water, and suggests that the ectoplasm may consist of layers of these substances which thus play an important rôle in absorption and secretion. Green and Jackson also consider that it exercises considerable influence on the transport of materials from cell to cell. This view of the lipoid nature of the plasmatic membrane is greatly supported by the work of Czapek* on the surface tension of the external limiting layer of the protoplasm.

Lipoids may of course represent an intermediate product between the fats and proteins, for it is a well-known fact that fats may develop in cheese, but according to Nierenstein† in

^{*} Czapek: "Ueber eine Methode zur direkten Bestimmung der Oberflächenspannung der Plasmahaut von Pflanzenzellen," Jena, 1912. + Nierenstein: "Proc. Roy. Soc., Lond.," B., 1911, 83, 301.

such cases the fats are derived not from the proteins, but from other substances, such as cholesterol.

Also the view has been put forward that they are the means of setting up the change in zymogens which leads to the formation of enzymes. More recently Palladin* has suggested that there is a relationship between lipoids and respiration, for the more of these substances extracted from seedlings the more was the respiration depressed. Possibly the lipoids, which contain phosphorus, act in a similar way as the phosphates in alcoholic fermentation.

* Palladin: "Ber. deut. bot. Gesells.," 1910, 28, 120; Palladin and Stanewitsch: "Biochem. Zeit.," 1910, 26, 351.

SECTION II.

THE CARBOHYDRATES.

UNDER the general heading of carbohydrates are included the sugars, starches, gums and celluloses, all of which substances play an important part in the economy of the plant. such as starch, sugars, inulin and glycogen, are all-important as food materials, and are stored up for future use in various Speaking generally, the carbohydrate reserves are at a maximum in the autumn and sink to a minimum in the early summer, after the expansion and growth of the leaves and young shoots. Other carbohydrates, such as the celluloses, play an important part in the mechanics of the plant, since they are largely concerned in the formation of cell-walls; some have merely a transitory existence, such as maltose; and finally, some possibly may be degradation products-e.g., mucilages. Notwithstanding the differences in the physiological significance of the various types of carbohydrates in the plant, these substances are all closely related chemically, being composed of the same elements—carbon, hydrogen and oxygen-united together in a similar fashion.

The term carbohydrate originated through the erroneous conception that these substances were compounds of carbon with water, since the proportion of hydrogen to oxygen in all of them is the same as in water, as may be seen from the formula for grape sugar, which is $C_6H_{12}O_6$, but which might be written C_66H_9O .

CLASSIFICATION OF CARBOHYDRATES.

On purely physical grounds such as appearance, solubility in water, taste, etc., the carbohydrates may be roughly divided into sugars and non-sugars; the systematic classification of the carbohydrates is, however, based upon their behaviour

towards hydrolytic agents, such as mineral acids or enzymes. Thus there are a considerable number of naturally occurring sugars containing five and six carbon atoms which cannot be hydrolysed; such sugars form a group known as monosaccharides.* On the other hand many sugars are known which on hydrolysis break up into two molecules of monosaccharide according to the equation

$$C_{12}H_{22}O_{11} + H_2O = 2C_6H_{12}O_6$$

Such sugars are known as disaccharides.

Similarly sugars which on hydrolysis give three molecules of monosaccharide as follows-

$$C_{18}H_{32}O_{16} + 2H_2O = 3C_6H_{12}O_6$$

are termed trisaccharides

Finally, carbohydrates, such as starch and cellulose, which on hydrolysis yield an unknown number of molecules of monosaccharides are classed as polysaccharides.

The nomenclature of the monosaccharides is based on the number of carbon atoms in their molecules, those containing five being called pentoses, while those containing six atoms are known as hexoses. For this reason the use of the terms monose and biose in place of monosaccharide and disaccharide is to be deprecated owing to the confusion which is liable to result therefrom.

$$I. \ Sugars. \begin{cases} & \text{Pentoses} \ (C_5H_{10}O_5). \ Arabinose, \ Xylose, \\ & \text{Rhamnose, Fucose, } \\ & \text{Quinovose.} \end{cases} \\ & \text{Hexoses} \ (C_6H_{12}O_6). \ Sorbose, \ Levulose, \\ & \text{Sorbose, Galactose, } \\ & \text{Mannose.} \end{cases} \\ & \text{Disaccharides} \ (C_{12}H_{22}O_{11}). \ Sucrose, \ Maltose, \ Isomaltose, \\ & \text{Cellobiose, Trehalose, Agavose, } \\ & \text{Lupeose, Mellobiose.} \end{cases} \\ & \text{Trisaccharides} \ (C_{18}H_{32}O_{16}). \ Raffinose, \ Melecitose, \ Stachyose.} \end{cases}$$

$$& \text{II. Non-sugars or Polysaccharides.} \end{cases} \\ & \text{Starches} \ (C_6H_{10}O_5)_{\text{n}}. \ \ \text{Starch, Dextrin, Mannoseanes, Galactosanes, } \\ & \text{Glycogen, Inulin.} \\ & \text{Glycogen, Inulin.} \\ & \text{Gums} \ \left(\begin{pmatrix} a \\ b \end{pmatrix} \ \text{Mucilages and Pectic bodies.} \right) \\ & \text{Celluloses} \ (C_6H_{10}O_5)_{\text{n}}. \end{cases}$$

^{*} The artificially prepared tetroses, heptoses, octoses, and nonoses also belong to this group, but as they do not occur in nature, as far as is known, they need not be considered here.

CONSTITUTION AND ISOMERISM OF SUGARS.

The analysis of any one of the hexose sugars, such as dextrose, levulose, galactose or mannose, would yield the same result; viz., 40 per cent of carbon, 6'ô per cent of hydrogen, and 53'3 per cent of oxygen; and this notwithstanding the fact that these sugars are different substances.

From the results of an analysis, it is possible to determine the simplest ratio of the atoms to each other in the molecule by dividing each percentage by the atomic weight of the corresponding element, and then determining the simplest numerical ratio between the resulting numbers:—

$$C = \frac{40^{\circ}0}{12} = 3^{\circ}3^{\circ}; H = \frac{6^{\circ}6}{1} = 6^{\circ}6^{\circ}; O = \frac{53^{\circ}3}{16} = 3^{\circ}3^{\circ}$$
$$\therefore C : H : O = 3^{\circ}3 : 6^{\circ}6 : 3^{\circ}3$$
$$= 1 : 2 : 1.$$

The formula CH_2O thus arrived at, is known as the Empirical Formula; it indicates the *ratio* of the number of different atoms in the molecule, but does not indicate their actual number. The formula which, while maintaining the above ratio, also shows the actual number of atoms present in the molecule, is known as the Molecular Formula; and it can only be assigned correctly when the molecular weight is known. Now the molecular weight of all these sugars is 180, hence their molecular formula must be $(CH_2O)_6$ or $C_6H_{12}O_6$.

Compounds such as the various hexoses which have the same molecular formula and yet are not identical are said to be isomers.

The carbohydrates exhibit two kinds of isomerism, known respectively as structural and stereo-isomerism.

Structural isomerism is well illustrated by the two sugars dextrose and levulose. A study of their reactions, which need not here be detailed, leads to the conclusion that they both contain five hydroxyl (OH) groups; that dextrose belongs to a class of compounds known as aldehydes, which are characterized by the group—CHO; and that levulose is a ketone and therefore contains the group = CO. These facts are all explained by the following constitutional formulæ:—

Levulose,
CH ₂ OH
Снон
Снон
Снон
co L
СНОН

Stereo-isomerism is the second type of isomerism, and is exhibited by the three sugars dextrose, mannose and galactose, all of which are aldehydes, and have therefore the same structural formula. The possibility of isomerism in this case is accounted for by the presence in these molecules of what are known as asymmetric carbon atoms. Writing the formula for dextrose once more in a slightly different way, it will be seen that the carbon atom printed in "clarendon" (C) has its four valencies attached respectively to the groups (CH₂OH.

Any carbon atom whose valency is satisfied by four different groups or elements, whatever their nature may be, is said to be asymmetric, since it is possible to represent it by either of two solid models which are not super-imposable, the one being the mirror image of the other; there exists, therefore, between two modifications of such an asymmetric carbon atom a difference due to the different spacial distribution of the four substituting groups around it. Now the isomerism existing between glucose and mannose is accounted for by their each containing one of the two possible modifications of this same asymmetric carbon atom. Similar considerations will show that each of the three carbon atoms marked with a star is also asymmetric, and it is therefore not surprising that it is possible to account for no less than sixteen different isomeric aldehyde sugars or aldoses; of these, however, relatively few have been found in nature.

The presence of an asymmetric carbon atom confers upon

a compound the property of optical activity, by which is meant the power of the substance to rotate to the right or to the left the plane of a beam of circularly polarized light passing through it. This phenomenon may be made use of for the estimation or identification of a sugar in solution (pp. 94, 95).

GENERAL REACTIONS OF SUGARS.

There are no reagents except that of Molisch (p. 58) which are of general application for the characterization of sugars, but there are two, namely phenylhydrazine and Fehling's solution, which react with by far the greater number of sugars and are consequently very largely used.

Phenylhydrazine, which was discovered by Fischer, reacts only with sugars containing either an aldehyde or ketone group to form, in the first place, phenylhydrazones, which in many cases are characteristic crystalline solids, but are usually soluble in water; this reaction may be illustrated thus:—

 $\label{eq:choose} \text{CH}_2\text{OH}(\text{CHOH})_4\text{CHO} + \text{H}_2\text{NNHC}_6\text{H}_5 = \text{CH}_2\text{OH}(\text{CHOH})_4\text{CH}: \text{NNHC}_6\text{H}_5 + \text{H}_2\text{O}} \\ \text{Dextrose or Glucose Phenylhydrazone}$

If, however, an excess of phenylhydrazine be employed, a second hydrazine complex is introduced into the compound, and the resulting substance is termed an osazone. Both glucose and levulose yield the same osazone,

 $\begin{array}{c} {\rm CH_2OH(CHOH)_3--C-CH:NNHC_6H_5} \\ {\rm N.NHC_6H_5} \end{array}$

which is called glucosazone.*

The osazones being, for the most part, insoluble in water, serve as a valuable means of isolating a sugar from a dilute solution; their identity can then be readily established by means of their crystalline form, melting point, solubility and optical activity.

A second very important reagent for sugars, depending for its utility, like phenylhydrazine, on the presence of the aldehyde or ketone group, is Fehling's solution. This substance, which is an alkaline solution of cupric oxide, acts upon a warm solution of a sugar as an oxidizing agent and, in parting with its oxygen, is converted into cuprous oxide; this reduction of

^{*}For details of the preparation of this substance, see under reactions for glucose, p. 62.

cupric oxide to cuprous oxide is accompanied by a visible change from a deep blue solution to a colourless one, with simultaneous deposition of the cuprous oxide as a reddish-brown precipitate. Any easily oxidized substance will thus reduce Fehling's solution, becoming itself oxidized; and, inasmuch as the aldehyde and ketone groups are readily oxidized, all sugars containing these groups will bring about this change. The reducing power of all sugars, however, is not the same, but it has been determined in most cases against a properly standardized Fehling's solution, and hence can be employed as a means of identifying or estimating the strength of a sugar solution. In view of what has been said, it will of course be seen that the experimental determination of the reducing power of a sugar is valueless if an unknown amount of any other easily oxidized substance is present in solution.

MONOSACCHARIDES.

A. PENTOSES.

The pentoses, which are sugars containing five carbon atoms, have the general formula $C_5H_{10}O_5$; they are not as a rule found free in plants, but may occur in a combined state. For example, in cherry or wood gum they occur in the form of pentosanes or pentosides, which substances may be regarded as anhydrides of pentoses, since they give rise to these sugars on hydrolysis in much the same way that starch on hydrolysis yields glucose.

GENERAL PROPERTIES OF PENTOSES.

- 1. They are not fermentable by yeast.
- 2. On distillation with hydrochloric or sulphuric acid, they are converted in furfural, which may be detected by its turning a solution of aniline acetate red.

$$C_5H_{10}O_5$$
—3 $H_2O = C_4H_3O.CHO$

This may easily be seen by boiling some wood shavings with concentrated hydrochloric acid in a test tube and allowing the escaping steam to impinge upon a piece of filter paper moistened with aniline acetate; * a pink colour is produced.

^{*} Prepared by mixing together equal parts of aniline, water and glacial acetic acid.

It should be noted that hexoses* will also produce this reaction, though to a much smaller extent, since the quantity of furfural produced from them is much less (not more than 0'2 per cent) than in the case of the pentoses. The chief product of the action of concentrated hydrochloric acid in hexoses is levulinic acid.

3. Warmed with concentrated hydrochloric acid (sp. gr. I'2) and a little orcinol, they produce a greenish-yellow colour which is soluble in amyl alcohol to a clear green solution having a characteristic absorption spectrum with bands between the C and D lines.

The reaction may be modified by adding a couple of drops of ferric chloride to the solution after it has been heated with hydrochloric acid and orcinol, when a bright green colour is produced.

N.B. This test is characteristic for pentoses.

4. By substituting phloroglucinol for orcinol in the above test, a red colour is produced, which changes to a brown precipitate; the latter is soluble in amyl alcohol, the solution having an absorption band between the D and E lines. This is the same reaction that is employed for the detection of lignified tissues; its use in this case depends on the fact that lignocellulose contains a pentose or furfural-yielding complex (see p. 145).

Dextrose and levulose when subjected to this test produce a yellow or brown colour.

5. Pentoses answer Molisch's test for carbohydrates. This test, which is also dependent on the formation of furfural from the sugar, consists in adding 2 c.c. of concentrated sulphuric acid to a mixture of the sugar solution with 2 drops of 15 per cent alcoholic solution of α -naphthol, which must be free from acetone. At the junction of the two liquids a green ring is produced, and over this a red zone; on cooling and shaking the colour changes to purple.

This test is given by all carbohydrates and glucosides, and proteins which contain a carbohydrate radicle.

- 6. They form osazones.
- 7. The pentoses reduce Fehling's solution.

^{*} Cf. Tollens: " J. f. Landw.," 1901, 39.

PROPERTIES OF INDIVIDUAL PENTOSES.

Arabinose.

Arabinose is best obtained by the hydrolysis of cherry gum with 4 per cent sulphuric acid; it can also be obtained by the hydrolysis of gum arabic and of peach gum. Arabinose has a very sweet taste, is dextro-rotatory, a_D in 10 per cent solution = $+105^{\circ}$, crystallizes in prisms, and melts at 160° ; it reduces Fehling's solution, and yields with diphenylhydrazine a characteristic diphenylhydrazone, melting at 218° .*

Xylose.

Xylose may be obtained by the hydrolysis of xylane or wood gum, and also from brewers' grains, maize fruits, straw, and various forms of cellulose. It is optically inactive, and crystallizes in prisms, melting at 144-145°. When oxidized with bromine it gives xylonic acid, which may be identified by the fact that it forms an insoluble double salt with cadmium bromide.†

Methyl Pentoses (CH₃C₅H₉O₅).

(a) Rhamnose.—Rhamnose, sometimes wrongly called isodulcite, has the empirical formula $C_0H_{12}O_5$, and is a pentose in which one of the hydrogen atoms has been replaced by a methyl group, its constitution being represented by the formula

сн₃снон снон снон снон сно

In common with other methyl pentoses it yields on distillation with hydrochloric acid, methylfurfural; this latter may be detected by warming a little of the distillate with an equal volume of concentrated hydrochloric acid, when a yellow colour is produced.

Rhamnose has been obtained by the hydrolysis of a number of glucosides, e.g., quercitrin, hesperidin, and xanthorhamnin, and also saponins. The substance forms glistening crystals, m.p. 93° ; $\alpha_D = +8.07^{\circ}$, and gives a phenylosazone melting at 180° .

^{*} Neuberg: "Ber. d. deut. chem. Ges.," 1900, 33, 2243. † Widstoe and Tollens: id., 1900, 33, 136.

- (b) Fucose.—Fucose, which is isomeric with rhamnose, may be obtained by the hydrolysis of sea-weeds by means of dilute sulphuric acid; it crystallizes in microscopic needles, and yields a hydrazone, m.p. 172-173°.
- (c) Quinovose, another methyl pentose isomeric with rhamnose, is produced by the hydrolysis of quinovite, a substance formed by boiling quinovin contained in the bark of Cascarilla hexandra with alcohol and hydrochloric acid.

B. HEXOSES.

There are no convenient general reactions for distinguishing hexoses from any other group of sugars, but each of the hexoses occurring in nature are readily identified by characteristic reactions.

GLUCOSE OR DEXTROSE.

The substance which is commonly known as grape sugar occurs, together with levulose or fruit sugar, in a number of sweet fruits, in honey, and in the seeds, leaves, roots, and blossoms of a great many of the higher plants; monosaccharides also obtain in lower plants; thus, Hunger describes them as occurring in small granules near the plastids in *Dictyota*. Glucose is formed by the hydrolysis of cane sugar, of glucosides, and of many polysaccharides, such as starch, cellulose, etc.

Preparation of Glucose.

The most convenient source for the preparation of glucose on a small scale is cane sugar. One hundred and twenty c.c. of 90 per cent alcohol mixed with 5 c.c. of fuming hydrochloric acid are heated at 45-50°; 40 grams of powdered cane sugar are now added, the mixture being kept thoroughly stirred. After two hours the solution is allowed to cool, and a little anhydrous glucose is added to induce crystallization. In the course of a few days the resulting crop of crystals is filtered off and washed with a little dilute alcohol; it is recrystallized by dissolving in half its weight of warm water and adding twice as much 90-95 per cent alcohol, filtering warm and setting aside to cool.

On a commercial scale glucose is best prepared by heating

freshly prepared potato or maize starch with dilute sulphuric acid in sealed copper vessels under 3 atmospheres pressure. When the hydrolysis is complete, the acid is removed as calcium sulphate by the addition of powdered chalk, and the filtered solution, after being decolorized by means of animal charcoal, is evaporated in a vacuum; a little anhydrous glucose in then introduced, and the syrup is allowed to crystallize at a temperature of 40°. Prepared in this way, the glucose forms a rather soft cake of small crystals; it is not a pure product, being contaminated with maltose, isomaltose (p. 73), and dextrin; it may, however, be purified by recrystallizing from aqueous alcohol.

Commercial dextrose is employed as a substitute for cane sugar for the sweetening of cheap jams, etc., but its sweetness is only about three-fifths that of cane sugar. The use of impure sulphuric acid containing arsenic for the hydrolysis of starch, and the subsequent employment of the glucose in the preparation of beer, has been the cause of the numerous deaths from arsenical poisoning.

Properties.

Glucose separates from alcoholic solution or from concentrated aqueous solutions at 30-35° in needle-shaped crystals, which are anhydrous; from cold aqueous solutions, however, it crystallizes with one molecule of water $(C_6H_{12}O_6 \ . \ H_2O)$ in the form of plates. It is readily soluble in water, but only very slightly soluble in absolute alcohol. It is readily fermented by yeast.

Glucose is dextro-rotatory, $a_{\rm p}=52^{\circ}5^{\circ}$; it is sometimes known as dextrose to distinguish it from the lævo-rotatory sugar levulose with which it is frequently found associated in ripe fruits.

Reactions.

I. In the presence of ammonia, glucose can reduce silver from its salts. A little glucose is added to a solution of silver nitrate to which have been added a few drops of caustic potash and just sufficient ammonia to redissolve the brown precipitate. On warming the mixture the silver is deposited on the sides of the test tube, forming a mirror.

2. Nylander's Test.—When boiled with a solution of glucose Nylander's reagent turns brown and finally black owing to the precipitation of bismuth oxide and metallic bismuth.

The reagent is prepared by dissolving 2 grams of bismuth oxynitrate and 4 grams of Rochelle salt in 100 grams of 10 per cent caustic soda solution.

- 3. Add to the solution basic lead acetate and ammonia. If glucose be present, a white precipitate comes down, which turns red. This reaction is not given by cane sugar.
- 4. Add to the solution a little copper sulphate solution and an excess of caustic potash. On warming, a yellow to red precipitate is formed. This reaction also is given by levulose and maltose, but not by saccharose.
- 5. On warming with Fehling's solution, a red precipitate is given by dextrose, levulose and maltose, but not by saccharose.
- 6. Add a little Barfoed's reagent and warm. A red precipitate floating as a thin film on the surface of the liquid indicates dextrose. This reaction is also given by levulose but not by cane sugar or maltose.

The reagent, which should be freshly made up, is prepared by dissolving 6.5 grams of copper acetate in 100 c.c. of water containing I gram of glacial acetic acid.

- 7. The addition to the solution of picric acid and caustic soda results in the formation of a blood-red coloration, due to picramic acid. This reaction is also given by other sugars.
- 8. On boiling the solution of glucose with an equal volume of caustic potash, a yellow-brown colour results; on acidifying with dilute nitric acid the colour lightens and a smell of burnt sugar is produced.
- 9. Glucose reacts with phenylhydrazine to give an osazone. To 5 c.c. of an approximately 5 per cent solution of glucose, add 4 or 5 drops of phenylhydrazine and about the same amount of glacial acetic acid. (If phenylhydrazine hydrochloride is used, add about enough solid to cover a threepenny piece and an equal quantity of sodium acetate.) Place the mixture in a boiling water bath for about half an hour and then remove; a golden yellow crystalline precipitate will have been formed. On examination under the microscope the needle-shaped crystals will be seen to be gathered together

in clusters resembling wheat sheaves. Glucosazone melts at 204-205° with decomposition; it is insoluble in water but soluble in alcohol, the solution being lævo-rotatory in contradistinction to that of maltose which is dextro-rotatory.

Microchemical Tests.

For microchemical tests for sugars, the reduction of copper salts in the presence of excess of alkali is generally employed, but these are not altogether satisfactory, owing to the amount of diffusion which takes place, and also because sucrose, if its presence in a tissue be suspected, must first be hydrolysed by boiling with acid before the reduction will take place.

Mangham* and others have obtained excellent results by the use of the osazone test for microscopic work; if properly performed, it is much more satisfactory than any other, and has the advantage of being a very delicate test for some sugars. For example, a '015 per cent solution of glucose will give a definite reaction. The main disadvantage of the method is in its comparative slowness.

Two solutions are required:

(a) I gram of phenylhydrazine hydrochloride dissolved in 10 grams of glycerine.

(b) I gram of sodium acetate dissolved in 10 grams of glycerine.

If necessary the solution of these substances may be hastened by means of heat, and before use the solutions should be filtered.

Glycerine is used because its penetrative power is greater than that of water, and also because it will not evaporate and deposit crystals of the substances used.

For use, one drop of each fluid is placed on a glass slip and mixed thoroughly. The section, which must be more than one cell in thickness, is laid in the mixture and covered with a cover glass. The preparation is heated on a hot water oven for about half an hour, and is then allowed to cool; the osazone crystals will form in varying degrees of rapidity.

In order that familiarity with the method may be gained, the reagents may be mixed on the slip with drops of sugar

^{*} Mangham: "New Phytol.," 1911, 10, 160; "Ann. Bot.," 1915, 29, 360.

solution of different concentrations heated for varying periods and examined periodically after cooling.

Maltose gives an osazone characterized by dense rosettes of lemon yellow crystals, which are broader and larger than those obtained with dextrose and levulose; the crystals may, however, take several weeks for their formation.

Dextrose and levulose may be distinguished by the fact that methylphenylhydrazine gives a crystalline osazone with levulose and not with dextrose. It is to be noted that the methylphenylhydrazine must be very pure.

LEVULOSE OR FRUCTOSE.

Levulose occurs in most sweet fruits and in honey, together with both cane sugar and dextrose, but usually in excess of the latter two. It is formed in equal quantity with dextrose by the hydrolysis of cane sugar, but being more strongly lævorotatory than dextrose is dextro-rotatory, the resulting mixture turns the plane of polarized light to the left, whereas the original cane sugar is dextro-rotatory; the resulting mixture is accordingly known as invert sugar and the process by which this change is produced is called inversion.

The separation of pure levulose from invert sugar on a small scale is not easy to carry out, but the operation is performed on a large scale by making use of the fact that on treating invert sugar with milk of lime the levulose is converted into an insoluble calcium compound, which may be filtered off and purified, while the glucose remains in solution. The easiest means of preparing levulose in the laboratory is to hydrolyse inulin by boiling I part of this substance with 5 parts of '5 per cent sulphuric acid* for one hour; the acid is then removed by means of barium carbonate, and the solution, after being treated with animal charcoal and filtered, is evaporated at a low temperature to a thin syrup. The latter is then crystallized from alcohol after sowing with a crystal of pure levulose. A modification of this method is employed for the manufacture of pure levulose. †

^{*} Düll ("Chem. Zeit.," 1895, 19, 216) recommends the use of oxalic acid; see also Wiechmann: "Z. d. Vereins Deut. Zuckerind.," 1891, 41, 331.

⁺ Cf. Stein: "Proc. Internat. Confer. Sugar Ind.," April, 1908.

Properties.

Levulose separates from alcohol in hard rhombic crystals, which have the composition $C_6H_{12}O_6$; from concentrated aqueous solutions, however, it crystallizes in needles with water of crystallization $2C_6H_{12}O_6$. H_2O . It is fairly soluble in hot absolute alcohol and ether, and may thus be separated from other sugars which are insoluble in these solvents. Levulose is strongly lavo-rotatory and exhibits slight muta-rotation; its rotatory power is very dependent on temperature, $a_p^{\ 20} = -93$ in a 10 per cent solution.

Reactions.

1. To a solution of levulose mixed with an equal volume of concentrated hydrochloric acid a few grains of resorcin are added. On warming, a deep red coloration results, and finally a brown-red precipitate. The precipitate is soluble in alcohol, giving a deep red solution.

This reaction is given by all keto-hexoses and by carbohydrates such as cane sugar and raffinose which give rise to them on hydrolysis.

- 2. Levulose gives the same reactions as dextrose with salts of copper and picric acid.
- 3. Levulose with milk of lime forms an insoluble compound; dextrose does not.
- 4. Levulose gives with phenylhydrazine the same osazone as glucose, namely glucosazone.
- 5. With methylphenylhydrazine it gives, in alcoholic solution, an osazone crystallizing in needles; m.p. 158°. (Distinction from glucose.)

SORBOSE.

Sorbose is a ketonic sugar produced by the fermentative oxidation of the alcohol sorbite contained in the sap of the mountain ash, *Pyrus Aucuparia*; this sugar probably does not exist as such in the plant, but is produced by oxidation as described.

GALACTOSE.

This sugar is formed as a product of the hydrolysis primarily of milk sugar, but also of the gums occurring in

peaches and plums, and from the so-called pectic substances occurring in carrots and pears; in all these cases it is accompanied by other sugars, which may be either hexoses or pentoses. Galactose also occurs in several plants belonging to the Caryophyllaceae. It is also formed by the hydrolysis of the trisaccharide raffinose, of the glucoside digitalin, and of a glucoside occurring in the ivy.

Preparation.

Galactose is best prepared by boiling milk sugar for six hours with four times its weight of 2 per cent sulphuric acid; the solution is then evaporated, and a few crystals of galactose are added to induce crystallization. After some time the crude galactose crystallizes out; it is purified by dissolving in four-fifths of its weight of water, and mixing the resulting solution with twice its volume of 93 per cent alcohol; the precipitate is filtered off and dried.

Properties.

Galactose crystallizes in minute hexagonal crystals, which melt at 168° . It is strongly dextro-rotatory, $a_{\rm D}=83.8^{\circ}$, and exhibits muta-rotation; it ferments completely, but rather more slowly than glucose.

Detection.

- 1. The hexagonal form of the crystals is characteristic of galactose.
- 2. It gives a phenylhydrazone (m.p. 158-160°) which is lævo-rotatory.
- 3. It reduces Fehling's solution somewhat more slowly than glucose; I c.c. Fehling's solution $\equiv 5^{\circ}11$ mg. galactose.
- 4. On oxidation with nitric acid it yields mucic acid. Five grams of substance are heated in a beaker with 6 c.c. of nitric acid (sp. gr. 1.15) until two-thirds of the liquid have been evaporated off. After twelve hours the mucic acid formed will have separated, and may be washed with 10 c.c. of water. If other insoluble substances, such as cellulose, etc., are present, place the filter paper with the solid in a dilute solution of ammonium carbonate to extract the mucic acid as ammonium

salt. Filter once more, and evaporate the filtrate almost to dryness, and acidify with nitric acid; the precipitate is pure mucic acid.

MANNOSE.

Mannose may be obtained by the hydrolysis of a form of mannane contained in salep mucilage (Orchis Morio) and from several other so-called hemi-celluloses contained in peas, coffee beans, date stones, etc. It is most conveniently prepared by the hydrolysis of the hemi-cellulose contained in ivory nuts, the fruits of Phytelephas macrocarpa; turnings from these seeds,* obtained in the manufacture of vegetable ivory buttons, are heated over a water bath for six hours with twice their weight of 6 per cent hydrochloric acid. After filtering, the solution is boiled with animal charcoal, neutralized and precipitated with phenylhydrazine acetate; the sugar may be isolated from the resulting phenylhydrazone by decomposing it with concentrated hydrochloric acid in the cold.

Mannose when dry is a hard crumbling substance, which, however, deliquesces and is readily soluble in water; it is only slightly soluble in hot alcohol and is insoluble in ether. It is dextro-rotatory, $[a]_0^{20} = + 14.36^\circ$ in 10 per cent solution, and is readily fermentable by yeast.

Detection

1. Mannose is most readily detected and estimated by means of its phenylhydrazone, which is almost insoluble in water, and forms almost at once on adding phenylhydrazine acetate to an aqueous solution of the sugar; the phenylhydrazone is soluble in a very large volume of boiling water, and separates in fine prisms from the solution on cooling. These crystals melt at 195-200°.

An excess of phenylhydrazine converts mannose into glucosazone, which is identical with the substance obtained under similar conditions from both glucose and levulose.

2. Mannose reduces Fehling's solution, 1 c.c. = 4.307 mg. mannose.

^{*} Fischer and Hirschberger: "Ber. deut. chem. Gesells.," 1889, 22, 3218.

Both mannose and galactose are sometimes considered to be transitory substances; an idea supported by the fact, observed by Hérissey,* that seminase is found in the seeds of lucerne; and that during germination cane sugar is relatively abundant, while mannose and galactose are not found, at any rate in any quantity.

DISACCHARIDES.

CANE SUGAR, SUCROSE OR SACCHAROSE.

Cane sugar is one of the most widely distributed substances to be found in the vegetable kingdom. Besides forming about 20 per cent of the juice of the sugar cane, Saccharum officinarum, and about 10 to 20 per cent of that of the beetroot, it is found in varying quantities in the wood of maple and birch, and in Sorghum saccharatum; it occurs, moreover, in wheat, maize, barley, in carrots and in madder root. In most sweet fruits it is found together with a greater or lesser quantity of dextrose and levulose, which may possibly have been formed from it by hydrolysis. It also is found in the leaves of many plants associated with glucose and maltose. The following table, compiled by Kulisch, gives the relative proportions of cane sugar and hexoses found in various fruits.

				Cane Sugar.	Hexoses
Pine apple				11.33	1.08
Strawberry				6.33	4.98
Apricot .		4		6.04	2.74
Ripe banana	ı.			5.00	10.00
Apple .				1-5.40	7-13:00

In honey practically only invert sugar † is found, although the sugar found in the flowers by the bees is commonly cane sugar. The hydrolytic agent in this case is, however, most probably the formic acid secreted by the bees.

The two chief sources for the preparation of cane sugar on a manufacturing scale are the sugar cane and the beet. The processes used in both cases are more or less similar, and consist in obtaining the juice, purifying it, concentrating it and, lastly, crystallizing it. The juice is generally obtained from the cane by crushing, as much as 85-95 per cent of the juice

^{*} Hérissey: "Rev. gen. Bot.," 1903, 15, 345, 369, 406, 444.

[†] A mixture of dextrose and levulose, see p. 64.

being expressed in this way; in some cases it is extracted by diffusion, which consists in immersing the cane in water, when the sugar diffuses out of the cells into the surrounding water while the indiffusible colloids remain behind. The crude juice is then boiled with milk of lime, in order to neutralize any acid present and to precipitate coagulable proteins, and is subsequently treated with sulphur dioxide. After filtering, the solution is concentrated in a vacuum and allowed to crystallize, the mother liquor being separated by centrifugalizing; the crystals may be used at once as brown sugar, or may be refined.

When the beet is used, the roots are first cut into slices and subjected to diffusion, the same quantity of water circulating through a series of vessels in such a manner that the fresh water first passes over material from which most of the sugar has already been extracted, and as the solution becomes more concentrated, it comes into contact with material which is increasingly richer in sugar. In this way the aqueous extract attains a concentration of from 12-15 per cent.* This solution is then boiled with lime and saturated with carbon dioxide to decompose any calcium saccharosate which may have been formed; it is then filtered and again saturated with carbon dioxide or a mixture of this gas and sulphur dioxide to precipitate the last traces of calcium, and also to decolorize it; the older process of filtration through animal charcoal is thereby rendered unnecessary; the solution is then boiled and filtered and the clear filtrate is concentrated in a vacuum and allowed to crystallize. The uncrystallizable residue which remains is known as molasses: a further yield of sugar may be obtained from this residue by the addition of lime to the cold solution or of strontia to the boiling solution whereby the cane sugar in the molasses is converted into the insoluble calcium or strontium saccharosate, which may be filtered off and decomposed by a current of carbon dioxide into cane sugar and calcium or strontium carbonate. The molasses are sometimes fermented for the manufacture of rum or may be used for cattle food: they are also used in the manufacture of boot blacking.

By suitable methods of cultivation and plentiful use of nitrogenous and potash fertilizers the amount of sugar con-

^{*} The residue remaining after the extraction of the sugar is employed for cattle food.

tained in the beet has been raised from 10.6 per cent in the period 1880-90 to about 15 per cent in the period 1900-10, and the beetroot is gradually displacing the sugar cane as a source of sucrose. Owing to exceptionally favourable weather conditions the yield in the year 1908-9 rose to about 18.5 per cent, but individual beets have been known to contain up to 27 per cent of sugar.

Properties.

Cane sugar crystallizes from water in monoclinic crystals which do not contain water of crystallization; it is readily soluble in water and only slightly soluble in alcohol; it is dextrorotatory, its specific rotation being $a_{\rm p} = +66.5$.

When heated to 160° it melts to a glassy mass known as barley sugar, which gradually becomes crystalline again; if heated to 190-200° it is converted into an uncrystallizable brown substance known as caramel, which is used for colouring beer and wine.

Reactions.

- 1. Solutions of cane sugar heated with concentrated hydrochloric acid turn reddish pink.
- 2. If warmed with concentrated hydrochloric acid and a few crystals of resorcin a deep red colour is produced owing to the liberation of levulose.
 - 3. Cane sugar does not react with phenylhydrazine.
 - 4. Cane sugar does not reduce Nylander's reagent.
- 5. Solutions of cane sugar do not reduce Fehling's solution until they have been inverted by boiling for a short time with a few drops of dilute sulphuric acid; if then made alkaline and boiled with Fehling's solution reduction ensues.

If a solution in water is boiled with a few drops of mineral acid, the sign of the optical activity of the solution changes from + to -. This change, which is known as *inversion*, is due to the fact that the mineral acid hydrolyses the cane sugar, converting it into equal molecular proportions of the two sugars dextrose and levulose,

$$C_{12}H_{22}O_{11} + H_2O = C_6H_{12}O_6 + C_6H_{12}O_6$$

and since the optical activity of levulose is greater than that of dextrose the resulting invert sugar is lævo-rotatory.

Numerous experiments have been carried out with a view to determining the conditions which bring about this inversion. Aqueous solutions of cane sugar, if kept for some time, gradually become inverted, the change being somewhat accelerated by prolonged boiling.

Similarly cane sugar solutions when heated with acids undergo inversion, the rate at which the change takes place being a measure of the strength, or better, the chemical affinity, of the acid. Extremely small quantities of acid suffice to effect the change in a boiling solution; thus 80 parts of cane sugar dissolved in 20 parts of water are completely hydrolysed by heating in boiling water for one hour with an amount of hydrochloric acid corresponding to 0.005 per cent of the weight of the sugar; within certain limits, however, the action is accelerated by increasing the concentration of the acid. however, the acid is too strong and the heating be continued too long, the solution is liable to darken and decompose. Moreover, prolonged action, even at temperatures of 10-15°, of concentrated acids was found by Wohl* and by Fischer † to produce exactly the opposite phenomenon, known as reversion, by which the simple molecules, more especially those of levulose, are made to condense together to form complex dextrinlike substances, as well as a disaccharide isomaltose,

6. Sucrose is not directly fermentable by pure yeast.

MALTOSE.

This disaccharide has not such a wide distribution in the plant as has cane sugar. It occurs in the cell sap of leaves and is formed, at any rate in part, by the action of diastase on the starch. Maltose is produced in quantity during the germination of barley and other grains by a similar enzyme action. The action is hydrolytic, and may be represented approximately by the following formulæ:—

The same change can also be brought about by the careful hydrolysis of starch with sulphuric acid.

^{*} Wohl: "Ber. deut. chem. Gesells.," 1890, 23, 2092. † Fischer: "Ber. deut. chem. Gesells.," 1890, 23, 3687.

Maltose is also formed by the action of diastase and other enzymes on glycogen.

In preparing maltose from starch, it is not necessary always to act on the starch contained in barley, potato-starch serving equally well, the diastase which is employed is usually introduced in the form of malt, which is barley that has been allowed to sprout and is then killed by suddenly heating to a temperature sufficient to stop the further growth of the barley without destroying the diastase. The malt is then stirred up with starch and water, and kept at a temperature of 60-62° for about half an hour; by the end of this time about 80 per cent of the starch has been converted into maltose and 20 per cent into dextrin. Dextrin itself is also converted into maltose by diastase, but the reaction is very slow, and in practice sufficient time is not allowed to effect this change.

Properties and Reactions.

Maltose is readily soluble in water, and crystallizes from this solvent in slender white needles, having the composition $C_{19}H_{92}O_{11}$, H_9O .

- 1. Maltose reduces Nylander's reagent, but not Barfoed's reagent.
- 2. Maltose reduces Fehling's solution without previous hydrolysis, and can therefore be estimated directly by this means.
- 3. When treated with phenylhydrazine, as described under glucose, it gives an osazone (m.p. 206°), which is soluble in seventy-five parts of boiling water, and can be crystallized from this solvent in rosettes of plates or broad needles resembling sword blades; alcoholic solutions of maltosazone are dextro-rotatory. (Distinction from glucosazone.)
- 4. On hydrolysis, by boiling with dilute sulphuric acid, maltose breaks up into two molecules of glucose:—

$$C_{12}H_{22}O_{11} + H_2O = 2C_6H_{12}O_6$$

the rotatory power of the solution being thereby diminished.

- 5. The aqueous solution is strongly dextro-rotatory: $a_p = + 137^\circ$; freshly-made solutions exhibit a higher rotation than older ones, owing to a negative muta-rotation.
 - 6. Unlike cane sugar, maltose is said to be directly fer-

mentable by yeast; this statement is, however, probably not strictly true, since pure cultures of yeast containing only zymase, the active enzyme which produces alcoholic fermentation, have no action on maltose. Ordinary brewers' yeast, however, contains maltase, which first hydrolyses maltose to grape sugar, which is then fermented by zymase. Only sugars containing six carbon atoms are fermentable by yeast (see p. 377).

ISO-MALTOSE.

Iso-maltose is a disaccharide which is isomeric with and closely related to ordinary maltose, its optical activity, $a_D = + 139 \cdot 140^\circ$, being almost the same as that of maltose; it is formed, together with ordinary maltose, when diastase acts on starch, provided the enzyme is not present in too large a quantity; the most favourable temperature for its production is 65-70°. Iso-maltose is also formed together with dextrin by the action of concentrated hydrochloric acid on glucose at a temperature of 10-15°, which accounts for the fact that it is not infrequently met with as an impurity in commercial glucose prepared by the action of hydrochloric acid on starch (see p. 61).

CELLOBIOSE.

Cellobiose is the name given to a disaccharide obtained by the hydrolysis of cellulose (see p. 135).

MYCOSE OR TREHALOSE.

Mycose or trehalose is the name given to a disaccharide found in various agarics, notably *Boletus edulis*, and also in moulds such as *Aspergillus niger*. It does not reduce Fehling's solution and is strongly dextro-rotatory, $a^{\rm p} = +199^{\circ}$. When boiled in acids it is slowly converted into glucose.*

AGAVOSE AND LUPEOSE.

Agavose † and lupeose ‡ similarly are disaccharides which have been isolated from the stem of *Agave americana* and lupin seeds respectively.

^{*} Winterstein: "Z. physiol. Chem.," 1894, 19, 70.

⁺ Michand and Tristan: "Amer. Chem J.," 1892, 14, 548.

[‡] Schulze: "Ber. deut. chem. Ges.," 1892, 25, 2213.

LACTOSE OR MILK SUGAR.

This disaccharide, though of considerable importance in the animal kingdom, is never found in plants and need not therefore be considered here.

TRISACCHARIDES.

RAFFINOSE.

This sugar occurs in cotton seeds, barley, eucalyptus, manna and also in the beetroot; the juice of this latter contains on an average about 15 per cent of cane sugar but only 0.02 per cent* of raffinose. The molasses from beet sugar refineries, however, contain from 2-3 per cent of raffinose (hence the name) and form the chief commercial source of this sugar.

As the concentration of the raffinose increases it tends to crystallize out together with the cane sugar in the form of mixed crystals having a peculiar and characteristic pointed appearance quite different from ordinary cane sugar.

Numerous methods † have been described for preparing pure raffinose from molasses, but as they are mostly rather tedious they will not be detailed here.

According to Bau,‡ raffinose may be extracted from cotton seeds by the following simple process. The powdered seeds, after being freed from fat by means of ether, are extracted with hot 70 per cent alcohol and the extract is heated with animal charcoal, filtered, and evaporated; on cooling raffinose crystallizes out and may be further purified by recrystallization from alcohol.

Raffinose crystallizes with 5 molecules of water in clusters of slender glistening needles or prisms whose composition is expressed by the formula $C_{18}H_{32}O_{16}$. $5H_2O$. It dissolves in water and in methyl alcohol, in which latter solvent cane sugar is only sparingly soluble, but is hardly soluble in ethyl alcohol, whereas cane sugar is appreciably soluble.

It is strongly dextro-rotatory, $aD = +104.4^{\circ}$, in 10 per cent

^{*}Strohmer: "Oest. Ung. Z. f. Zuckerind. u. Landw.," 1910, 39, 649.

[†]v. Lippmann: "Die Chemie d. Zuckerarten," 3rd ed., Braunschweig, Vol. II. p. 1628.

[‡] Bau : "Chem. Zeit.," 1894, 18, 1796.

solution, and consequently cane sugar in which raffinose occurs as an impurity appears to contain more than 100 per cent of sucrose when estimated polarimetrically; hence raffinose is sometimes known as "plus sugar".

It does not reduce Fehling's solution, nor does it react with phenylhydrazine.

On careful hydrolysis raffinose breaks up at first into levulose and a disaccharide—melibiose.

$$\begin{array}{ccc} C_{18}H_{32}O_{16} + H_2O = C_6H_{12}O_6 + C_{12}H_{22}O_{11} \\ Raffinose & Levulose & Melibiose \end{array}$$

On heating further the melibiose itself is broken up as follows:—

$$\begin{array}{l} C_{12}H_{22}O_{11}\,+\,H_{2}O\,=\,C_{6}H_{12}O_{6}\,+\,C_{6}H_{12}O_{6}\\ \text{Melibiose} & \text{Dextrose} & \text{Galactose} \end{array}$$

If boiled with mineral acid, therefore, raffinose gives rise to a mixture of dextrose, levulose and galactose.

According to Neuberg,* raffinose is hydrolysed by emulsin into cane sugar and galactose. (See below.)

Raffinose, unlike cane sugar, is completely fermented by bottom fermentation yeast to alcohol and carbon dioxide, whereas top fermentation yeast is only able to ferment it partially, converting the levulose complex into carbon dioxide and alcohol and leaving melibiose unattacked. These facts have been made use of by Bau† for detecting and for estimating raffinose.

Detection.

There are no rapidly performed characteristic tests for raffinose.

The only really reliable method of identifying it is to isolate the substance by precipitating the strontium compound in alcoholic solution, filtering off the precipitate and decomposing it by a current of carbon dioxide. The resulting solution is then evaporated and the residue extracted with alcohol to remove sucrose and other sugars which are more soluble in alcohol than raffinose. The pure substance should be identified by its crystalline form and optical properties.

^{*} Neuberg: "Bioch. Zeitschr.," 1907, 3, 519.

⁺ Bau: "Chem. Zeit.," 1894, 18, 1797; 1897, 21, 185; 1902, 26, 69.

Another way of identifying raffinose* is to add to the solution a little decoction of fresh yeast, to act as nutriment, and then to sterilize the solution; a pure culture of top fermentation yeast is then added to the solution and the fermentation is allowed to proceed in a thermostat at 31°; when it is completed, the solution is boiled with animal charcoal, filtered, and evaporated to a syrup: the latter is then, while still hot, poured into hot alcohol and on cooling it is filtered; the filtrate is then precipitated by mixing with 11 vols. of ether. After 24 hours the supernatant liquid is poured off and the residual syrup, which consists of melibiose, is converted into its osazone which is characterized by its crystalline form and melting point 178-9° †.

Finally, Neuberg t has proposed making use of emulsin for the identification of raffinose.

MELECITOSE.

This is a sugar which occurs in the sap of Larix europaea and in Persian manna; it crystallizes with two molecules of water in rhombic prisms, and is dextro-rotatory ($a_p = + 83^{\circ}$). It does not reduce Fehling's solution, and on hydrolysis yields first a molecule of glucose and a disaccharide-turanose, C₁₉H₂₉O₁₁—which subsequently itself breaks up into two molecules of glucose.

STACHYOSE.

$$C_{18}H_{32}O_{16}$$
, $_3H_2O$ or $C_{36}H_{62}O_{31}$, $_7H_2O$.

This substance may be obtained from the tubers of Stachys tuberifera. It forms plate-like crystals, which dissolve readily in water to give a faintly sweet solution, which is dextro-rotatory $(a_n = + 148^\circ)$. When boiled with dilute mineral acid it vields glucose, levulose, and galactose.§

SUGARS OF UNKNOWN MOLECULAR WEIGHT OR SUGAR-LIKE POLYSACCHARIDES.

So long as the molecular weight of a sugar is known, it is possible to classify it as a particular kind of saccharide, but

^{*} Bau: "Chem. Zeit.," 1897, 21, 185. † Ibid., 1902, 26, 69.

[†] Neuberg: "Bioch. Zeitsch.," 1907, 3, 519 and 535. § Planta and Schulze: "Ber. deut. chem. Ges.," 1891, 24, 2705.

when it is not known, the substance has to be classified, for want of a better name, as a sugar-like polysaccharide.

Gentianose and Lactosin are two such sugars, obtained respectively from the roots of Gentiana lutea and of Silene vulgaris. Both are dextro-rotatory, but the former on hydrolysis yields a lævo-rotatory mixture, while the latter yields a mixture containing 50 per cent of galactose.

ESTIMATION OF SUGARS.

A. VOLUMETRIC METHODS.

I. ESTIMATION BY MEANS OF FEHLING'S SOLUTION.

The principle of this method lies in the fact that certain sugars are capable of reducing copper sulphate in hot alkaline solutions to cuprous oxide, the presence of which is indicated by a yellow-red precipitate.

Fehling's solution is made up in two solutions:—

- A, containing 69:28 grams of pure crystallized copper sulphate in one litre of distilled water.
- B, containing 350 grams of Rochelle salt and 100 grams of caustic soda in one litre of distilled water.

The solution A must be made up very accurately, whereas the quantities required for solution B need only be roughly weighed.

For use, 5 c.c. of A are mixed with 5 c.c. of B; the mixture is a deep blue colour, and is known as Fehling's solution. If correctly compounded, 10 c.c. of the solution contain '11 gram of cupric oxide, which is able to oxidize '05 gram of glucose.

This value is sufficiently correct for general purposes; it is, however, an approximation, and varies for different sugars, the factor for levulose, for instance, is '05144, whilst that for invert sugar is '04941. If it be desired to obtain very accurate results, it is better to standardize the solution by titrating 10 c.c. with a solution of glucose of known strength. Such a solution may be obtained by dissolving '95 gram of pure crystallized cane sugar in 500 c.c. of distilled water and boiling for fifteen to twenty minutes with 2 c.c. of concentrated hydro-

chloric acid. The solution must then be neutralized by the addition of solid sodium carbonate, and made up to 1 litre; 50 c.c. of this solution contain '05 gram of glucose, and should reduce exactly 10 c.c. of Fehling's solution.

It frequently happens in titrating liquid extracts from plants, etc., that the cuprous oxide will not settle down, but remains suspended in the solution as a fine turbidity. In such cases the addition of a few drops of aluminium sulphate may sometimes cause the precipitate to subside; if not, it will be necessary to boil a fresh portion of the original solution and then to add lead acetate; after filtering, the filtrate is saturated with hydrogen sulphide to remove excess of lead, and the titration is then carried out on the filtrate after boiling off the hydrogen sulphide.

Plant extracts may also contain tannins which must be removed before estimating the sugars. There are various ways of doing this; and in all cases it is best to try the separation with a small portion of the material first, in order to determine the efficiency of the method.

- I. The aqueous solution is mixed with about half its volume of ethyl acetate, and thoroughly shaken. The mixture is then placed in a separating funnel until the two solvents have separated, when the lower aqueous part, containing the sugar, is drawn off, and with it the process is repeated. The aqueous solution is now warmed on a water bath until all the ethyl acetate is driven off, and the cooled solution tested for tannin.
- 2. To every 100 c.c. of the solution add 4 grams of pure lead carbonate. Shake thoroughly at intervals for about four hours. Test the filtrate for tannins.
- The chief objection to this method is that lead salts soluble in water may be formed.
- 3. Add gelatine solution until no more precipitate is formed. Filter and wash the precipitate very thoroughly. The filtrates resulting from any of these operations may then be examined for the sugar.

Estimation of Pentoses.

When pentoses alone are present, they may be estimated in the same way as glucose; if, however, they are mixed with other carbohydrates, or are present in the form of pentosanes other methods must be employed (see p. 86).

Estimation of Glucose.

The following precautions must be taken in estimating glucose by this and similar methods involving the reduction of metallic salts.

- 1. Any substances such as tannins which may have the power of reducing the salts used in titration must be removed.
- 2. The strength of the sugar solution must be weak, because the reducing power of sugar varies with the concentration, hence it is best to titrate a solution of about the same strength as that used for the standardizing of the Fehling's solution. This necessitates preliminary estimation; should the strength of the solution be much above this point, add a known volume of water until the strength approximates '5 per cent.

The titration, which should be completed as rapidly as possible in order to avoid reoxidation of the solution by the air, is performed as follows:—

Five c.c. of each of the solutions A and B are placed in a white porcelain basin and 40 c.c. of water added; the mixture is then boiled. The sugar solution is placed in a burette and is run into the hot copper solution about 3 c.c. at a time; after each addition the solution is boiled and the precipitate allowed to settle before the next addition is made. When the blue colour has disappeared, the amount of sugar solution used is noted.

A second titration is then carried out, and all the sugar required, less I c.c., to effect complete reduction, is run in at once; should this prove too small an amount of sugar, more is added drop by drop until decolorization results. The process is repeated until two readings are obtained which do not differ one from the other by more than 0.2 c.c., the one being a little too high and the other a little too low; the mean of these gives the correct result.

The chief difficulty in the titration lies in the detection of the end point; this may be ascertained by allowing the precipitate to settle, and then tilting the basin so as to view the clear liquid against the white of the dish. But if the observer's colour-sense be not very critical, an error is easily made, hence various methods have been suggested to determine accurately the end point.

- I. Filter off a small quantity of the solution, acidify it with acetic acid and add a little potassium ferrocyanide; the presence of unreduced copper is indicated by the formation of a brown coloration or precipitate of copper ferrocyanide.
- 2. Ling's reagent consists of I gram of ferrous ammonium sulphate and 1.5 gram of ammonium sulphocyanide dissolved in a mixture of 10 c.c. water and 2.5 grams of strong hydrochloric acid. The solution is decolorized immediately before use by adding a few pieces of granulated zinc. A dozen drops of the reagent are placed separately on a glazed white porcelain plate and a drop of the titration mixture is, from time to time, added to one of the drops; when no pink colour is produced, the titration is complete.
- 3. Harrison's indicator is made by adding a little starch paste to 100 c.c. of 10 per cent solution of potassium iodide; as this solution will not keep more than a few hours, it must be freshly prepared. One c.c. of the indicator is acidified by the addition of 10 drops of acetic acid and a little of the titration mixture is added. The presence of unreduced copper is indicated by the appearance of a red or blue colour; the absence of any colour marks the end of the reaction.

EXAMPLE.—Amount of sugar solution required to decolorize 10 c.c. of Fehling's:—

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      II'7 c.c.
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Now since 10 c.c. of Fehling's are completely reduced by '05 gram of glucose,

```
    ∴ 11.6 c.c. of the solution contained .05 gram glucose.
    ∴ 100 c.c. , , , (05 × 100 / 11.6 ) ...
    = 4.31 per cent.
```

Estimation of Galactose and Mannose.

The procedure is exactly the same as for glucose:-

1 c.c. Fehling's = '0511 gram galactose '= 4307 gram mannose.

Estimation of Cane Sugar.

Cane sugar does not reduce Fehling's solution; it is therefore necessary to invert it in order to make the estimation. To do this, take a known volume of the sugar solution and add a sufficiency of strong hydrochloric acid to make it about a 10 per cent solution of the acid; heat on a water bath for about a quarter of an hour, at 70° C. Then neutralize with sodium carbonate, make up to a known volume and titrate.

The inversion of cane sugar may be represented thus:-

$$C_{12}H_{22}O_{11} + H_2O = C_6H_{12}O_6 + C_6H_{12}O_6$$

The molecular weight of cane sugar is 342, and the amount of invert sugar this will give on inversion is, from the equation, 360. In other words, I gram of glucose corresponds to $\frac{342}{342}$ = '95 gram of cane sugar. The titration result must therefore be multiplied by '95.

Estimation of Maltose.

Three points must here be remembered: firstly, that maltose will reduce Fehling's solution; secondly, that this reduction may not be complete, and therefore the maltose must be inverted before it is titrated; thirdly, that the reducing power of maltose is not the same as glucose, I gram of maltose having the same reducing power as '62 gram of glucose. From the equation representing the inversion of maltose, it may be found that I gram of maltose gives I'05 gram of glucose; and, as I gram of maltose has the same reducing power as '62 gram of glucose, it follows that I gram of maltose after inversion gives an increased reducing power, viz.:—

1'05 - '62 = '43 gram glucose,
.: '43 gram glucose = 1 gram maltose,
and 1 gram glucose =
$$\frac{1}{43}$$
 gram maltose,
=2'32 grams maltose.

The titration result, which represents glucose, must therefore be multiplied by 2.32.

Estimation of Mixtures of Sugars.

In many cases it is possible to isolate the different sugars in solution, and estimate them separately by means of Fehling's solution or by some other method, and this separation must be accomplished when their action on Fehling's solution is similar. For example, it may be desired to estimate the amount of levulose and dextrose in a solution. Add to the dilute solution some ammoniacal lead acetate; both sugars are precipitated as lead compounds. Filter off and wash the precipitate; suspend the precipitate in water and pass through it a current of carbon dioxide. The lead compound of glucose alone is decomposed, and the glucose goes into solution. Filter off and thoroughly wash the levulose lead compound, and then suspend it in water and decompose it with sulphuretted hydrogen.

Similarly, should these two sugars be mixed with cane sugar, the latter, on the addition of ammoniacal lead acetate, remains in solution, and thus is easily separated.

Inasmuch as this method is somewhat tiresome, the following methods may be followed whenever possible:—

GLUCOSE AND SUCROSE.

- I. Take 100 c.c. of the mixture and titrate with Fehling's solution.
- 2. Invert 100 c.c. of the mixture by the method given, and titrate.

The first operation gives the amount of glucose = a.

The second operation gives the original amount of glucose together with that due to the inversion of the cane sugar = b.

$$(b-a) \times 95 = \text{sucrose}$$
.

GLUCOSE AND MALTOSE.

Proceed exactly as for glucose and sucrose:-

a = amount of sugar before inversion. b = amount of sugar after inversion.

From the reasons already given under maltose, it follows that—

 $(b-a) \times 2.32 = \text{maltose},$ and $a - (\text{maltose} \times .62) = \text{glucose}.$

CANE SUGAR AND MALTOSE.

Cane sugar is inverted by citric acid, while maltose is not; this fact may be made use of in the estimation:—

 Add to 100 c.c. of the solution 5 grams of crystallized citric acid, and heat on the water bath for about one hour. Neutralize and titrate.

Reducing power = a.

2. Completely invert another 100 c.c. of the solution with hydrochloric acid; neutralize and titrate.

Reducing power = b; then $(b-a) \times 2.32 = \text{maltose}$, and $(a-\text{maltose} \times .62) = \text{sucrose}$.

GLUCOSE, CANE SUGAR, AND MALTOSE.

 Take 100 c.c. of the solution and titrate. The result includes the glucose together with maltose.

Reducing power = a.

Take another 100 c.c. of the solution, invert with citric acid, and then titrate. The result includes the glucose, and the invert sugar obtained from the cane sugar, together with maltose.

Reducing power = b.

3. Take a final 100 c.c. of the solution, and completely invert with hydrochloric acid. The result represents the whole of the sugars.

Reducing power = c.

Following the same reasoning as before:-

 $(b-a) \times "95 = \text{cane sugar},$ $(c-b) \times 2 \cdot 32 = \text{maltose},$ and $a - (\text{maltose} \times "62) = \text{glucose}.$

II. ESTIMATION BY MEANS OF PAVY'S SOLUTION.

The chief disadvantage connected with the use of Fehling's solution in the estimation of glucose is the difficulty in observing the end point of the titration owing to the red precipitate of cuprous oxide; this may be overcome by using Pavy's solution, which contains ammonia which dissolves the cuprous oxide with the formation of a colourless solution. As before, two solutions are necessary.

- A. 8:316 grams of pure crystallized copper sulphate are carefully weighed and dissolved in one litre of distilled water.
- B. 40.8 grams Rochelle salt. 40.8 grams caustic potash. 600 c.c. strong ammonia (1880). Distilled water to one litre.

In making up the mixture B great accuracy is not essential. For titration 25 c.c. of A (very accurately measured) are mixed with 25 c.c. of B. The complete reduction of 50 c.c. of Pavy's solution is effected by '025 gram of glucose.

Pavy's solution may also be prepared from Fehling's solution as follows: 120 c.c. of Fehling's are mixed with 300 c.c. of strong ammonia (*880) and 400 c.c. of 12 per cent potash solution. The mixture is then made up with distilled water to one litre.

Method.—Fit a 250 c.c. flask with a well-fitting cork bored



with two holes, one to contain an outlet tube and the other the nozzle Pour into the flask of the burette. 50 c.c. of Pavy's solution and 50 c.c. of distilled water; mix thoroughly and introduce a little powdered glass. Dilute the sugar solution with a 10 per cent solution of ammonia, in order that 50 c.c. shall be about equivalent to 50 c.c. of the Pavy solution. Bring the Pavy solution to the boil by means of a small flame, and run in the sugar solution I c.c. at a time. Having thus roughly ascertained the amount of sugar required, accurate readings are to be obtained by running in nearly all the requisite sugar at once, and then drop by drop until the end point is reached.

The following precautions are very important:-

 The operation must be carried out rapidly, else all the ammonia is driven off and the cuprous oxide is precipitated. 2. The Pavy solution must be boiling throughout the titration, else air will enter the flask, owing to the lowered temperature, and the solution of cuprous oxide will be oxidized

III. ESTIMATION BY MEANS OF BENEDICT'S SOLUTION.

In this method the difficulty of the red precipitate of cuprous oxide obscuring the end point is overcome by carrying out the reduction in the presence of potassium thiocyanate whereby the cuprous oxide is converted into an insoluble white compound, and thus the disappearance of the last trace of blue colour from the solution is easy to observe.

The solution is prepared as follows:-

200 grams sodium citrate,

200 grams crystallized sodium carbonate or 75 grams of the anhydrous salt,

125 grams potassium thiocyanate

are dissolved in water, made up roughly to 800 c,c., and filtered.

Eighteen grams of pure crystallized copper sulphate dissolved in 100 c.c. of water are poured slowly with constant stirring into the above solution. Five c.c. of a 5 per cent solution of potassium ferrocyanide are now added as a further precaution against the formation of cuprous oxide, and the whole is then carefully made up to 1000 c.c.

The above solution, which will keep indefinitely without any special precautions, is of such a strength that

25 c.c. = 0.05 gram glucose.

The titration is performed as follows:-

Twenty-five c.c. of Benedict's solution are placed in a 4 oz. flask with 3 or 4 grams of anhydrous sodium carbonate and a few lumps of broken porcelain to prevent bumping; the mixture is kept boiling vigorously while the sugar solution is run in until the blue colour just disappears. The sugar solution may be run in rapidly at first, but towards the end it should be run in drop by drop.

The volume of solution run in contains the equivalent of 0.05 gram glucose from which the strength may be calculated.

This method is easier to work with than Fehling's solution, and gives very accurate results.

B. GRAVIMETRIC METHODS.

Estimation of Pentoses.

These compounds may, according to Neuberg,* be estimated by conversion into their diphenylhydrazones,

$$C_5H_{10}O_4NN(C_6H_5)_2$$

and subsequent weighing; this method is, however, not always suitable.

The ease with which furfural can be produced from pentoses has led to the following method of estimation, which is due to Kröber†:—

A weighed quantity of substance ‡ (usually about 5 grams) is placed in a 300 c.c. flask provided with a cork bored with two holes, through one of which passes a tap-funnel, and through the other a splash preventor, such as is used in a Kjeldahl distillation. Through the tap-funnel 100 c.c. of hydrochloric acid (sp. gr. 1.06, containing about 12 per cent HCl) are then added, and the contents of the flask are distilled briskly; when 30 c.c. have passed over, the distillation is interrupted and the contents of the receiver are poured into a beaker with a 400 c.c. graduation mark; a fresh quantity of 30 c.c. of hydrochloric acid (sp. gr. 1.06) is now added through the tap-funnel, and the distillation is continued until 30 c.c. more have distilled over; the new distillate is again transferred to the beaker, 30 c.c. more acid are added to the flask, and the whole process is repeated; altogether about a dozen distillations, each lasting ten minutes, are required to carry over the last traces of furfural. In order to ascertain whether the distillate still contains furfural, a drop of the liquid is placed on a

^{*} Neuberg: "Ber. deut. chem. Gesells.," 1900, 35, 2243; see also Maurenbrecher and Tollens: "Ber. deut. chem. Gesells.," 1906, 39, 3578.

⁺ Kröber: "J. Landw.," 1900, 48, 357, and 1901, 49, 7.

[‡] The amount chosen should be sufficient to produce from '03 to '03 gram of phloroglucide,

filter paper next to a drop of aniline acetate solution; * if no red colour appears when the two liquids come in contact with each other, the solution is free from furfural, and the distillation can be discontinued.

The furfural contained in the united distillates is then precipitated from solution by means of phloroglucinol which reacts according to the equation:—

$$C_5H_4O_2 + C_6H_6O_3 = C_{11}H_6O_3 + 2H_2O$$

To this end about the amount of phloroglucinol † likely to be required by the furfural obtained is dissolved in hydrochloric acid (sp. gr. 1°06), and added to the furfural solution, and the total volume is then made up to 400 c.c. with more of the same acid. The solution at once turns yellow, then becomes turbid, and, on the next day, the greenish-black precipitate of the phloroglucide is filtered off on to a tared Gooch crucible; the precipitate is washed with 150 c.c. of water, dried for four hours at 97°, then cooled in a desiccator and weighed in a weighing bottle. From the weight (a) of the precipitate, which under ordinary conditions should lie between 003 and 0°3 gram, the weight of furfural, pentose or pentosane may be calculated by substituting the value of (a) in one of the following formula:—

a lies between 0.03 and 0.3 gram. Furfural = $(a + .0052) \times .5185$ Pentose = $(a + .0052) \times 1.0075$ Pentosane = $(a + .0052) \times .8866$

in which '0052 is the weight of phloroglucide, which remains in solution under the conditions of the experiment as given above.

If the precipitate weighs less than 0.03 gram or more than 0.3 gram, one of the following formulæ must be employed:—

* This is best prepared, according to Tollens, by shaking up equal volumes of aniline and water in a test tube and adding glacial acetic acid drop by drop until the turbid solution suddenly becomes clear.

†The phloroglucinol employed must be pure. To ascertain this, test as follows: Dissolve a small quantity in a few drops of acetic anhydride, heat almost to boiling and add a few drops of concentrated sulphuric acid; a violet colour indicates the presence of diresorcinol; if more than a faint coloration appears, the sample should be rejected.

‡This is necessary to prevent the phloroglucide, which is hygroscopic, from absorbing moisture.

```
Weight of precipitate < '03 gram. Weight of precipitate > 0'3 gram.

Furfural = (a + 0.0052) \times 0.517 Furfural = (a + 0.0052) \times 0.518

Pentose = (a + 0.0052) \times 0.0052 Pentosane = (a + 0.0052) \times 0.0052 Pentosane = (a + 0.0052) \times 0.0052 Pentosane = (a + 0.0052) \times 0.0052
```

According to Böddener and Tollens * a considerable saving in time may be effected by precipitating the phloroglucide in hot solution, i.e. between 80 and 85°. The reaction then takes place according to the equation—

$$C_5H_4O_2 + C_8H_6O_3 = C_{11}H_4O_2 + 3H_2O$$

so that the precipitate actually weighs less than the one produced in the cold; the precipitation is, however, complete in from one and a half to two hours. The weight of furfural corresponding to the precipitate so obtained may be calculated by adding 'OOI (to allow for the phloroglucide remaining in solution) and multiplying the resulting figure by 0.571. The number so obtained if multiplied by 1.703 gives the corresponding amount of pentose or if multiplied by 1.703 gives the amount of pentosane. The method is, however, not suitable if it is desired to estimate the methyl-pentosanes as distinct from the pentosanes, in which case Kröber's method as modified by Ellett † and Mayer † should be employed.

Estimation of Glucose.

Several methods \(\) have been devised for employing Fehling's solution for the gravimetric estimation of sugars, each method only giving accurate results if careful attention is given to all details of manipulation.

According to Allihn two solutions are required:-

A. Prepared by dissolving 34.6 grams of pure crystallized copper sulphate in water and making up the volume to 500 c.c.

B. Containing 173 grams of Rochelle salt and 125 grams of potassium hydroxide dissolved in water and made up to 500 c.c.

Thirty c.c. of each of the two solutions A and B are mixed together in a 300 c.c. beaker, diluted with 60 c.c. of water and

```
* Böddener and Tollens: "J. Landw.," 1910, 58, 232.

+ Ellett: "J. Landw.," 1905, 53, 13.

‡ Mayer: "J. Landw.," 1907, 55, 261.

§ Soxhlet: "J. f. prakt. Chem.," 1880, [2], 21, 227; Allihn: id., 1880, [2], 23, 63; Pflüger: "Pflüger's Archiv," 1898, 69, 399.
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heated to boiling; 25 c.c. of the sugar solution, so prepared as not to contain more than 0.25 gram of glucose, are then added, and the boiling is continued for exactly two minutes; * at the end of this time the supernatant liquid should be still blue; if not, the sugar solution was too strong and a fresh experiment must be started using a more dilute sugar solution. The liquid is then filtered through a weighed asbestos Gooch crucible which has been previously washed first with water and then successively with 10 c.c. of alcohol and a like quantity of ether, and has been dried for half an hour in a steam oven before weighing. The precipitate is then similarly washed with hot water and finally with 10 c.c. of alcohol and 10 c.c. of ether, and is dried for half an hour in a steam oven. After cooling in a desiccator, the Gooch crucible is weighed again.

The weight of cuprous oxide multiplied by the factor o 8883 gives the weight of copper, from which the amount of dextrose may be calculated by reference to the table on pp. 87-88.

An alternative method consists in mixing as before 30 c.c. of each of the solutions A and B, diluting them with 60 c.c. of water and heating by immersing the mixture in a boiling water bath for six minutes; 25 c.c. of the sugar solution, containing not more than 0.25 gram of glucose, are then boiled and added to the copper solution; the mixture is then heated in the boiling water bath for another ten minutes. The precipitated cuprous oxide is thereupon filtered, as before, on a tared asbestos Gooch crucible, washed, dried, and weighed.

As the results obtained by different workers are liable to vary somewhat, it is best for each worker to determine for himself what results he obtains when using a glucose solution of known strength; by dividing the weight of glucose known to be in the solution by the weight of cuprous oxide obtained, a factor is found which on subsequent occasions can be used for multiplying into the weight of cuprous oxide obtained, in order to give the corresponding amount of glucose.

A sufficiently accurate result can, however, usually be obtained by employing the factors given in the following table for

^{*} According to Pflüger this is not sufficiently long.

							
Copper,	Grape Sugar.	Соррег.	Grape Sugar.	Copper.	Grape Sugar,	Copper.	Grape Sugar.
5	Sug	ος	Sug	တိ	Sug	တိ	Sug
Mgram.	Mgram.	Mgram.	Mgram.	Mgram.	Mgram.	Mgram.	Mgram.
10	6.1	68	34.8	126	64.3	184	94.5
11	6•6	69	35'3	127	64.7	185	94.7
12	7.1	70	35.8	128	65.5	186	95.2
13	7·6 8·1	71 70	36·3 36·8	129	65.7 66.2	187 188	95'7
14	8.6	72 73	37.3	130 131	66.2	189	96·3
16	9.0	74	37.8	132	67.2	190	97'3
17	9.5	75	38.3	133	67.7	191	97.8
18	10.0	76	38.8	134	68.2	192	98.4
19	10.2	77	39.3	135	68.8	193	98.9
20 21	11.0	78 70	39·8 40·3	136	69.3 69.8	194	99'4
22	11.2	79 80	40.8	137	70*3	195 196	100.2
23	12.2	8 1	41.3	139	70.8	197	101.0
24	13.0	82	41.8	140	71.3	198	101.2
25	13.2	83	42.3	141	71.8	199	102.0
26	14.0	84	42.8	142	72.3	200	102.6
27 28	14.2 15.0	85 86	43°4 43°9	143 144	72 · 9	201 202	103.1
29	15.2	87	43 9	145	73.9	203	104.5
30	16.0	88	44.9	146	74.4	204	104.2
31	16.2	89	45*4	147	74'9	205	105.3
32	17.0	90	45.9	148	75.2	206	105.8
33	18.0	91	46.4	149	76 ° 0	207 208	106.3
34 35	18.5	92 93	46°9 47°4	150 151	76·5	200	100'8
36	18.0	93	47.9	152	77.5	210	107.9
37	19.4	95	48.4	153	78.1	211	108.4
37 38	19.9	96	48.9	154	78.6	212	100.0
39	20.4	97	49*4	155	79'1	213	109.2
40 41	20 . 0	98	49'9	156 157	79.6 80.1	214	110.0
42	21.0	99 100	50°4	158	80.7	216	111.1
43	22.4	IOI	51.4	159	81.5	217	111.6
44	22.9	102	51.9	160	81.4	218	112.1
45	23*4	103	52.4	161	82.2	219	112.7
46	23'9	104	52.0	162	82.7	220	113.5
47 48	24 . 4	105 106	53°5	163 164	83 · 3	22I 222	114.3
49	25.4	107	54.2	165	84.3	223	114.8
50	25.9	108	55.0	166	84·3 84·8	224	115.3
51	26.4	109	55'5	167	85*3	225	115.9
52	26.9	110	56.0	168	85.9	226	116.4
53 54	27.4 27.9	III II2	56·5	169 170	86•4 86•q	227 228	117.4
55	28.4	112	57.5	171	87.4	220	118.0
56	28.8	114	58•0	172	87.9	230	118.2
57	29.3	115	58.6	173	88.5	231	119.0
58	29.8	116	. 59'1	174	89.0	232	119.6
59 60	30°3	117	60.1	175 176	89·5	233 234	120°1 120°7
61	31.3	110	60.6	170	90.2	234 235	121.5
62	31.8	120	61.1	178	91.1	236	121.7
63	32*3	121	61.6	179	91.6	237	122.3
64	32.8	122	62.1	18o	92.1	238	122.8
65 66	33.3	123	62.6	181	92.6	239	123.4
67	33·8 34·3	124 125	63 · 7	182 183	93.1 93.1	240 241	123.0
/	343	-43	~3 /	13	93 /		44

5	ن به	ı,	ن به ا	Copper.	ن و	1 5	ے ہ
i d	Grape Sugar.	ă	gan	ă.	Grape Sugar.	ă	gan
Copper.	Sg	Copper.	Grape Sugar.	Š	5 g	Copper.	Grape Sugar.
Mgram.	Mgram.	Mgram.	Mgram.	Mgram.	Mgram.	Mgram.	Mgram.
		g.u		g.u	176111111	g.u	
242	125'0	298	155.4	354	186.6	410	218.4
		299	156.0	355	187.2	411	210,3
243	125°5	300	156.5	355 356		412	219.9
244 245	126.6	301	157.1	357	188.3	413	220'4
		302		357 358	188.0	414	221.0
246	127'1		157.6				221.0
247	127.6	303		359	189.4	415	
248	128'1	304	158.7	36o	190.0	416	222'2
249	128.7	305	159'3	361	190.6	417	222'8
250	129'2	306	159.8	362	191.1	418	223.3
251	129.7	307	160.4	363	191.7	419	223.9
252	130.3	308	160.0	364	192.3	420	224.5
253	130.8	309	161.2	365	192.9	421	225'1
254	131.4	310	162.0	366	193'4	422	225.7
255	131.0	311	162.6	367	194.0	423	226.3
256	132.4	312	163.1	368	194.6	424	226.9
257	133.0	313	163.7	369	195.1	425	227.5
258	133.2	314	164.2	370	195'7	426	228.0
259	134.1	315	164.8	371	196.3	427	228.6
260	134.6	316	165.3	372	196.8	428	229'2
261	135.1	317	165.9	373	197'4	429	229.8
262	135.7	318	166.4	374	198.0	430	230.4
263	136.2	319	167·0	375	198.6	431	231.0
264	136.8	320	167.5	376	199.1	432	231.6
265	137'3	321	168.1	377	199'7	433	232.2
266	137.8	322	168.6	378	200.3	434	232.8
267	138.4	323	169.2	379	200.8	435	233'4
268	138.0	324	169.7	38o	201'4	436	233.9
26g	139.5	325	170'3	381	202'0	437	234'5
270	140.0	326	170'9	382	202'5	438	235.1
271	140.6	327	171.4	383	203'I	439	235*7
272	141.1	328	172.0	384	203.7	440	236.3
273	141.7	329	172.5	385	204.3	441	236.0
274	142.2	330	173.1	386	204.8	442	237.5
275	142.8	331	173.7	387	205 4	443	238.1
276	143.3	332	174.2	388	206.0	444	238.7
277	143'9	333	174.8	389	206.2	445	239'3
278	144'4	334	175'3	390	207'I	446	239.8
279	145.0	335	175.9	391	207.7	447	240'4
280	145.2	336	176.2	392	208.3	448	241.0
281	146.1	337	177.0	393	208.8	449	241.6
282	146.6	338	177.6	393	209'4	450	242.2
283	147.2	339	178.1	394	210'0	451	242.8
284	147.7	339	178.7	395 396	210.6	452	243'4
285	148.3		179'3	397	211.5	452	243 4 244'0
286	148.8	341	179.8	397 398	211.7		244 6
287	140.0	342	180.4		213.3	454 455	244 0
288		343	180.0	399	212.0	455	
	149'9	344	181.2	400 401			245.7
289	150.2	345	182.1		213.2	457 458	246.3
290	151.0	346	182.6	402	214.1		246.9
291	151.6	347		403	214.6	459	247.5
292	152.1	348	183.2	404	215.2	460	248.1
293	152.7	349	183.7	405	215.8	461	248.7
294	153.2	350	184.3	406	216'4	462	249'3
295	153.8	351	184.9	407	217.0	463	249'9
296	154.3	352	185.4	408	217.5		
297	154'9	353	186 · o	409	218.1		
		1			1		

converting a given weight of cuprous oxide into glucose, cane sugar (after inversion), lactose, or maltose:—

		Glucose, Levulose, or Galactose.	Cane sugar.	Lactose.	Maltose.
Cuprous oxide		.5042	*4790	·68 ₄₃	.8132
Cupric oxide		*4535	.4308	·6153	.7314
Copper .		•5634	. 5395	. 7709	•9089

Supposing the weight of cuprous oxide obtained by the oxidation of a lactose solution to be 185, then the weight of lactose corresponding to this would be—

The chief source of error in this method lies in the possibility of the cuprous oxide containing impurities, such as silica or alumina, derived from the alkali, in which case, of course, its weight would be too high; to overcome this objection several methods have been devised, such as reducing the cuprous oxide to metallic copper, or depositing the copper electrolytically and weighing this; or else oxidizing the cuprous oxide to cupric oxide and weighing again.

The factors for converting metallic copper and cupric oxide into sugars are also given in the above table.

Estimation of Glucose as Osazone.

The following method of estimating glucose as osazone in the products of the action of malt upon starch is recommended by Davis and Ling:* Twenty c.c. of solution containing 2-3 grams of starch products per 100 c.c. are mixed with 1 c.c. of phenylhydrazine and 1.5 c.c. of 50 per cent acetic acid. After heating for an hour † over a water bath, the liquid, which has by this time evaporated to a small bulk, is filtered through a tared Gooch crucible, and the crystalline osazone is washed with 20-30 c.c. of boiling water, so that the total filtrate does not exceed 50 c.c.; the precipitate is then dried in a steam oven and weighed; under these conditions, 0.1 gram of glucose gives 0.0505 gram of glucosazone.

^{*} Davis and Ling: "Journ. Chem. Soc., Lond.," 1904, 85, 24.

⁺ The heating should not be continued for more than one hour.

C. POLARIMETRIC METHODS,

The polarimeter is much used in ascertaining the strength of sugar solutions, but before describing the mode of using it, it is desirable to consider briefly the principles which are involved.

When a ray of light enters a crystal of any system other than the cubical, it is broken up into two rays, the ordinary and the extra-ordinary, provided the beam of light is not coincident with the optical axis of the crystal. This phenomenon is known as double refraction.

These two rays, the ordinary and the extra-ordinary, do not behave similarly; the former conforms to the ordinary laws of refraction, but the latter does not; further, the two rays are polarized in directions at right angles to one another.

In order to make use of these facts, it is necessary to be able to examine the extra-ordinary ray alone; that is, the two rays must be separated one from the other. This is effected by a Nicol's prism, which consists of two plates of Iceland spar fixed together by means of Canada balsam. A ray of light enters one side of the prism, and is broken up into the ordinary and the extra-ordinary ray; on reaching the layer of balsam, the former is totally reflected, whilst the latter passes on through the other plate and emerges at the side opposite to its entry. If a second Nicol be placed in the path of this ray, the latter will pass through in different amounts according to the angle which the second prism makes with the first. the interposed Nicol be parallel to the first Nicol, the ray will pass through entirely; if the second Nicol be rotated, the light passing through will be less and less in amount until, when the two prisms are at right angles to each other, no light passes at all. If the rotation be continued, the light will again pass through in gradually increasing quantities until the prism has been rotated through an angle of 180° from its original position, when the whole light will again pass through freely.

Many liquids and solutions of solids possess what is known as optical activity, which means that they can rotate the plane of vibration of a ray of polarized light passing through them; so that, on emergence from the liquid, the new plane is inclined either to the right or to the left of the original plane.

This is known as the rotation of the plane of polarized light.

Laurent's Half-Shadow Polarimeter.—This apparatus consists of a tube containing two Nicol's prisms, of which one is fixed and is known as the polarizer, while the other can be rotated and is called the analyser. A quartz plate which covers half the field of vision is fixed just behind the polarizer.

The liquid or solution to be examined is contained within a glass tube with polished ends, and is placed in position between the quartz plate and the analyser. The analyser is fixed in a tube which can be rotated, the degree of rotation being read from a divided circle. Leaving out of consideration the quartz plate, the beam of polarized light passes through the liquid and so becomes rotated; it follows, therefore, that the vibration plane of the analyser will no longer be at right angles to the plane of polarization of the light striking it, therefore light will enter the analyser, and in order to bring about complete extinction, the analyser must be rotated either to the right or to the left. This angle of rotation is a measure of the optical activity of the substance under observation, and according to the direction of rotation, the substance is termed dextro- or lævo-rotatory. In Laurent's polarimeter the illumination is obtained from a sodium flame, and this light before entering the tube containing the liquid must pass through the plate of quartz. When the instrument is set in the zero position, the whole field is equally illuminated, but on introducing the liquid, one half of the field becomes the darker; equal illumination can be obtained by rotating the analyser. If this position be passed, the field is once more unequally illuminated, but in a reverse manner, that is to say, the half which was originally dark is now light, and vice versa.

As the exact position of equal illumination is somewhat difficult to determine, several readings should be made and the mean of these taken as the correct value.

The specific rotation of a substance is defined as the angular rotation produced by a column of liquid I dm. in length, which contains I gram of the active substance in each cubic centimetre. This quantity is expressed by the symbol $a_{\rm po}^{\rm 20}$, the numeral indicating the temperature at which the measurements were made, and the letter D standing for the yellow

line of the sodium flame which is used as the source of illumination. The use of this quantity a_D for determining the number of grams of active substance in a given solution will be rendered apparent from the following considerations.

Supposing we have a solution containing an unknown number of grams, m, of active substance per c.c., and we fill a tube of length / dm.* with this solution and observe its angular rotation to be a.

If a layer 1 dm. long containing 1 gram
$$\left\{\begin{array}{lll} \text{ of substance in 1 c.c. of liquid produces a rotation} \end{array}\right\} a_D$$

Then ,, ,, l ,, ,, ,, m ,, ,, ,, $m = m + l$, $m = l$, $m = m + l$, $m = l$, $m = m + l$, $m = l$

The angle of rotation is determined as follows:--

- I. Find the zero reading when no solution is between the polarizer and analyser. For this purpose the mean of at least three readings, differing by only two or three minutes, should be taken.
- 2. Fill the tube with the liquid, taking care to avoid the introduction of air-bubbles.
- 3. Insert the tube and determine the new reading at which the illumination of both halves of the field is equal. The mean of three readings should again be taken.

The difference between the initial and the final readings is the angle of rotation.

The following experiment performed on a solution known to contain glucose may be quoted in illustration of the method:—

Initial reading of polariscope, without any solution = 0° 30′ Final ,, ,, , with glucose ,, = 3° 45′ Difference (a) = 3° 15′ or
$$3^{\circ}25^{\circ}$$
 Length of tube containing the solution (t) = 2 dms. Specific rotation of glucose $(a_{20}^{\circ}) = 52^{\circ}5^{\circ}$

From which
$$m = \frac{3.25}{2 \times 52.5} = .0309$$

... the strength of the solution is 3.00 per cent.

^{*} The length of the tube must be expressed in decimetres.

It is of course obvious that correct values can only be obtained by this method on the assumption that the liquid contains only a single optically active substance.

Some substances, e.g. glucose, exhibit the phenomenon of muta-rotation, that is to say, the rotation of their solutions varies according to the length of time that they have been made up; the maximum rotation is given by a freshly-made solution, but the rotatory power gradually decreases until it finally becomes steady. The attainment of the final condition is greatly accelerated by warming the solution in the presence of a little alkali, but the solution must of course be cooled before a reading is taken.

POLYSACCHARIDES.

The second great group of carbohydrates, namely the nonsugars or polysaccharides, are substances of high molecular weight, mostly amorphous and insoluble in water. Like the di- and tri-saccharides, the polysaccharides on hydrolysis break up into sugars containing five or six carbon atoms, and they may therefore be looked upon as anhydrides of these substances.

In the absence of any exact knowledge regarding their molecular weights, their formulæ are written $(C_0H_{10}O_5)_n$ or $(C_5H_8O_4)_n$ according as they give rise to hexoses or pentoses on hydrolysis. The value of "n" has not been determined as yet for any particular case, but there is reason to believe that it is fairly high. The various methods adopted for the elucidation of this point have led to such widely different results that a description of them here would not serve any useful purpose.

CLASSIFICATION.

The polysaccharides may be classified as follows:-

- I. Starches and Dextrins, including Glycogen, Inulin, Mannane and Galactane ($C_6H_{10}O_5$)_n.
- II. Gums, which comprise (a) Natural Gums and Pentosanes; (b) Mucilages and Pectic Bodies.
 - III. Celluloses (C₆H₁₀O₅)_{n.}

STARCHES.

The general formula for all substances belonging to this group is $(C_6H_{10}O_5)_m$, which indicates that on hydrolysis they yield hexoses; for this reason they may be termed hexosanes. The hexoses produced however are different, and the group may therefore be subdivided as under, the basis of the classification being the nature of the sugar.

Starches or Hexosanes.—Ordinary starch, Dextrin, Glycogen, Lichenin, Paradextrane, etc.
Levulosanes.—Inulin, Phlein, Graminin, Triticin, Synanthrin, etc.
Mannosanes.—Mannane, Paramannane, Mannocellulose.
Galactosanes.—Galactane, Paragalactane.

DEXTROSANES.

Starch or Amylum.

Starch is one of the most widely distributed substances in the vegetable kingdom; it may be found in green leaves as a temporary reserve of the photosynthetic products; as a more or less permanent reserve food-material it occurs in seeds and fruits, where it is not infrequently accompanied by other reserves, for instance proteins; in the vegetative parts, such as tubers, the living cells of the pith, medullary rays, and cortex of roots and stems; and also in the latex of certain plants, e.g. Euphorbia. When especially stored, the amount of starch may be considerable; thus in cereals it may form from 50 to 70 per cent of the dry weight of the grains, and in potatoes from 15 to 30 per cent of the dry weight of the tubers. As is well known, starch grains from different sources show much variety in size and shape, and occur in association with plastids, in which, as Schimper demonstrated, they have their origin.

Many monocotyledonous plants are characterized by the absence of starch, for example Scilla nutans, Phleum pratense, Allium, etc., but in some of these cases starch granules may occur in the guard-cells of the stomates, in the bundle sheath of the leaves, and also in the bulbs at the base of the growing shoots; further, in certain plants which normally form sugar, e.g., Musa, Hemerocallis, and Muscari, starch will appear when much sugar accumulates. On the other hand, many

members of this same class of plants are fairly constant starch producers, e.g., Lilium tigrinum, Pontederia cordata, Ananas sativa, Canna indica, Tra lescantia virginica, Juncus communis, and Alisma Plantago. There are many peculiarities in this occurrence of starch in the Monocotyledons; for instance, in Scilla nutans it is absent, whilst in Scilla siberica it is quite abundant; further, the former plant, if fed with cane sugar in a solution of suitable strength, does not form it, while, on the other hand, starch-free plants of Scilla siberica under the same treatment do form starch, the experiment, of course, being carried out in the absence of light. In the plant starch occurs, as is well known, in the form of variously shaped microscopic bodies composed of concentric layers; the granules have an organized structure and possess the power of double refraction.

Preparation of Starch.

The method of preparation varies according to the source employed. From wheat flour it may be obtained by stirring up this material thoroughly with water, and allowing the mixture to stand until the gluten contained in the flour undergoes fermentation, when it dissolves and may be removed by washing. On a small scale the separation is most conveniently effected by kneading some flour in a muslin bag which is held under a stream of water. The starch granules are hereby washed through the muslin, while the gluten remains behind in the bag as a sticky grey mass.

Starch may also be obtained from potatoes by macerating them with water and separating the non-starchy material from the starch by filtration. The starch is then allowed to settle at the bottom of the water, when it is collected and dried.

Purification.

Malfitano and Moschkoff * give the following method for the purification of starch: A one per cent colloidal solution of starch is frozen and then allowed to melt. When melted, most of the starch is deposited in a floccular precipitate, whilst the clear liquid contains some starch and the greater part of the

^{*} Malfitano and Moschkoff: "Compt. rend.," 1910, 151, 817.

mineral impurities. On repeating the operation four or five times, the purified product yields less than '02 per cent of ash.

Properties.

Air-dried starch contains a considerable quantity of water, as much as 20 per cent being not uncommon; it can be made to part with this water by carefully heating to 100°. If heated to about 200° it is converted into a sticky soluble substance, which is probably a mixture of isomeric substances of the empirical formula $C_6H_{10}O_5$, known as British gum or dextrin (q.v.).

Starch is quite insoluble in cold water, but if heated with water the granules swell and burst, a slimy opalescent mass known as starch paste being formed. The consistency of this paste varies of course with the concentration, and also with the particular kind of starch employed; this may be accounted for by assuming that some starches are richer than others in the constituent which produces the viscosity (p. 100). If a dilute starch paste be filtered, a gelatinous residue remains on the filter paper; the filtrate contains some starch, since it gives a blue colour with iodine, but it is doubtful whether the liquid is a true solution; it is more likely a colloidal solution in which the particles are sufficiently small to pass through the pores of the filter paper.

With regard to the chemical nature of starch granules there are considerable differences of opinion. The researches of Nägeli have shown that when starch is treated with dilute hydrochloric acid, malt extract, or saliva, a considerable portion goes into solution, leaving a transparent skeleton undissolved. The soluble portion, which gives a blue colour with iodine, Nägeli regarded as the true starch constituent of the granule, and named it granulose; on the other hand, the undissolved skeleton, which is not turned blue by iodine, he considered to be of a cellulose nature, and called it starch cellulose or amylocellulose.

On the other hand, Meyer * was of opinion that starch granules consisted essentially of two substances known respectively as α and β amylose. The former, which was insoluble,

^{*} Meyer: "Unters. u. d. Stärkekörner," Jena, 1895.

he regarded as an anhydride which could be converted into the soluble β variety by the action of superheated steam.

He also thought that when starch is acted upon by hydrochloric acid it is converted into amylo-dextrin, and considered that amylo-cellulose, which Nägeli regarded as an original constituent of the starch granule, was in reality identical with amylo-dextrin, and therefore a secondary product of the action of acid on the amylose.

According to the view of recent workers, notably Maquenne and Roux,* and Fernbach and Wolff,† starch granules consist of two substances: amylo-cellulose, or amylose, as they describe it later, and amylo-pectin. The term amylo-cellulose is, however, employed in a different sense from that assigned to it by Nägeli. It is, according to these authors, the principal constituent, and is partly soluble in boiling water and completely soluble in water under pressure; in solution it gives a blue colour with iodine, and is converted into maltose by malt, but in the solid state it is not acted upon by these reagents. soluble form is produced by heating the insoluble one with water under pressure, and the insoluble form may be recovered from the solution by cooling, a process which is known as "reversion". The insoluble amylo-cellulose is probably identical with the substance described by Nägeli under that name, in that it is not coloured by iodine; it is, however, not regarded as differing essentially from the soluble form (Nägeli's granulose), but rather as being a polymer of it or a different crystalline variety.

The second constituent, amylo-pectin, is a mucilaginous substance of an entirely different nature, which is not coloured blue by iodine and dissolves in malt extract, without, however, being converted into a sugar; it swells up without dissolving when heated with water. According to Maquenne ‡ and Roux, it is amylo-pectin which produces the gelatinization of starch in the form of starch paste, which substance may therefore be regarded as a colloidal solution of amylo-cellulose (amylose) thickened by an insoluble gelatinized slimy material,

^{*} Maquenne and Roux: "Compt. rend.," 1903, 137, 88; 1905, 140, 1303.

⁺ Fernbach and Wolff: id., 1904, 138, 819.

[†] Maquenne: "Bull. Soc. Chim.," Paris, 1906, [3], 35, 1, and "Ann. Chim. Phys.," 1904, [8], 2, 109; Maquenne and Roux: "Ann. Chim. Phys.," 1906, [8], 9, 179.

the amylo-pectin. Amylo-pectin, moreover, tends to retard the reversion of soluble amylo-cellulose into the insoluble form, and hence there is a quantity of the soluble form present in the starch granule which is able to dissolve in boiling water; when, however, the amylo-pectin is removed, the pure insoluble amylocellulose or amylose (as the authors prefer to call it) is produced.

A new form of soluble starch has recently been described by Fernbach.* It is obtained by pouring a 1 or 2 per cent aqueous suspension of potato starch into a large excess of pure acetone and shaking vigorously; a flocculent precipitate is thus obtained, which, when filtered and ground up in a mortar with more acetone and then dried in a vacuum, yields a light white powder which is completely soluble in cold water. The aqueous solution passes through filter paper and yields a very pure blue colour with iodine.

Brief mention may be made of the ideas held regarding the physical nature of starch grains. As is well known, the granules not infrequently exhibit a more or less well-marked stratification which years ago was thought to correspond to the alternation of day and night.

The "apposition" theory held that new layers were added to those already formed, each layer being separated from the next by a thin film of air. Nägeli came to the conclusion that the lamellation was due to the differences in the water-content of the several layers, and that the grain was made up of minute particles, the micellæ, which are of the prismatic order, surrounded by a film of water and embedded in a matrix. growth of the grain took place by a process of intussusception, that is to say, new material was intercalated between the micellæ, and either gave rise to new micellæ, or was used up in increasing the size of the old ones. Schimper expressed the idea that the grains were really of a sphæro-crystalline nature, which view was modified by Meyer, who says that the grain is made up of two kinds of needle-shaped crystals composed respectively of α and β amylose; he also states that in those grains which are coloured red with iodine, for example those found in the cells of the root-cap of Allium Cepa, in the seedcoats of Chelidonium and in Oryza sativa, var. glutinosa, dextrin

^{*} Fernbach: "Compt. rend.," 1912, 155, 617.

and amylo-dextrin occur. On the other hand, the ordinary grains which are coloured blue with iodine, are made up almost entirely of sphæro-crystals of amylose arranged in layers.

According to Kraemer,* the starch grains of the potato are composed of colloid and crystalloid substances arranged in lamellæ which are distinct and separate one from the other. At the point of origin of growth, the hilum, and in the alternate lamellæ, the colloid preponderates and is associated with the crystalloid cellulose; in the other lamellæ the crystalloid granulose is in the greater proportion. He also states that the peripheral layer is elastic and porous, and may be an anhydride of cellulose. Dennison also has expressed the view that the outer layer of the grain is different from the more internal parts, and may be a carbohydrate not fully polymerized to starch.

The view that both crystalloid and colloid materials occur in the starch grain is held by many; in addition to those referred to above, Czapek may be mentioned; it is, however, not universally held, Fischer,† who believes that the grain is composed of colloid substances, being a dissentient.

Action of Acids on Starch.

The action of acids on starch varies according to the strength of the acid, the duration of the action, and the temperature of the experiment. To complicate matters, there are considerable divergences in regard to the interpretation of the results obtained by the different workers. As an illustration of the very different effects which may be produced under different conditions, the following experiments may be quoted.

By acting on starch at the ordinary temperature with 12 per cent commercial hydrochloric acid for twenty-four hours, Brown and Morris found that granules, while retaining their external features, had acquired the power of dissolving in hot water without the formation of paste. The addition of alcohol to such a solution caused the immediate precipitation of a white substance known as soluble starch, which is turned blue by iodine, is strongly dextro-rotatory, $[a]_D = 202^\circ$, and

^{*} Kraemer: "Bot. Gaz.," 1902, 34. + Fischer: "Beih. bot. Centr.," 1905, 181.

does not reduce Fehling's solution. On the other hand, if starch is boiled for some time with dilute hydrochloric acid, it is converted into glucose, a fact which is made use of in estimating starch.

That maltose is also produced as an intermediate product of the acid hydrolysis of starch has been shown by Fernbach and Schoen,* and also by Weber and Macpherson,† who have proved it to be present in commercial glucose (see p. 60). Accompanying the conversion of starch into glucose there is, however, the formation of varying quantities of gummy substances known as dextrins (q.v.); it is, however, not known for certain whether these dextrins are formed directly by the action of the acid on the starch, or whether they are produced by the condensing action of the acid on the glucose already formed; there is moreover great difference of opinion with regard to the nature and number of these substances which are formed.

Action of Malt or Diastase on Starch.

The action of malt extract or diastase (see p. 368), like that of mineral acids, is primarily a hydrolytic one in which the starch is converted into a sugar; but the diastase does not carry the hydrolysis so far as the acid, the sugar produced being a disaccharide, maltose, and not a monosaccharide. At the same time some dextrin is also formed. Although the mechanism of this reaction is very complex and has led to a great deal of discussion, the ultimate change may be conveniently represented by the following formulæ:—

Action of Bacteria on Starch.

By the action of *Bacillus materans* on 5 per cent starch paste, Schardinger \ddagger has obtained two crystalline substances which he describes as α and β dextrin. Pringsheim and Langhans \S ascribe to these compounds the formulæ $(C_eH_{10}O_5)_4$ and $(C_6H_{10}O_5)_6$ respectively; they have obtained

^{*} Fernbach and Schoen: "Bull. Soc. Chim.," 1912, [iv.], 11, 303.

⁺ Weber and Macpherson: "J. Amer. Chem. Soc.," 1895, 17, 312.

[‡] Schardinger: "Zeitsch. f. Natur. u. Genussm.," 1903, 6, 874.

[§] Pringsheim and Langhans: "Ber. deut. chem. Ges.," 1912, 45, 2533.

from the former a crystalline disaccharide $(C_6H_{10}O_5)_2$, and from the latter a crystalline trisaccharide $(C_6H_{10}O_5)_3$. All these four compounds have a sweet taste and, according to the authors, they are representatives of a new class of crystalline polysaccharides which they term amyloses.* The substances are accordingly named di-, tri-, tetra-, and hexa-amylose.

Reactions.

- The appearance of the grains under the microscope and their action on polarized light in the presence of water are well known.
- 2. The most characteristic reaction of starch is the blue colour produced with iodine. The composition of this blue substance varies; it contains, on an average, about 18 per cent iodine, and cannot be formed unless a small quantity of hydriodic acid, which is always present in small amounts in ordinary solutions of iodine, be present. The blue colour is discharged on heating the solution, but reappears on cooling. The dried substance may, however, be heated to 100° without undergoing alteration. It is stated that those parts of the grain which are particularly rich in granulose are the most affected by the iodine.

If the starch grains are very small, or relatively so few in number that they might easily be overlooked, Meyer's procedure may be followed. A section of the material to be examined is cut, and is first treated with a fairly dilute solution of iodine in potassium iodide, the excess of the reagent is then removed, and the section is irrigated with a concentrated aqueous solution of chloral hydrate. This causes the starch grains to swell, and at the same time the other cell contents are dissolved, as are also the starch grains in time.

- 3. For microscopic work, the action of dilute aqueous solutions of gentian violet and of safranin is sometimes useful, as they stain the colloidal parts more deeply.
- 4. Starch grains are insoluble in cold water, but in hot water they swell up and form an opalescent solution which, if strong enough, will on cooling eventually form a paste.
- * The choice of this term is unfortunate in view of the various uses to which it has already been put by other authors, such as Meyer, Maquenne and Roux, etc. (see pp. 99, 100).

- 5. Starch is precipitated from its aqueous solution by alcohol or by basic lead acetate (cf. Inulin and Dextrin).
 - 6. Potash causes the grains to swell and finally to dissolve.
- 7. Boil a little starch paste solution with a few drops of dilute sulphuric acid in a test tube, and from time to time remove a little of the solution, cool it and test with iodine solution; when the starch has been converted into dextrin the blue colour at first formed will give way to a plum colour. If boiled too long only dextrose will remain which gives no colour with iodine. The solution will, however, after making alkaline, reduce Fehling's solution.
- 8. Cautiously heat a little starch on a porcelain basin until it has acquired a light fawn colour. Cool and extract with cold water, and filter; the dextrin produced being soluble in cold water is thus separated from the starch. On adding iodine to the solution a plum colour is produced.

Estimation of Starch.

A colorimetric method of estimating starch, depending on the depth of the blue colour produced with iodine, has been described by Dennstedt and Voigtländer.*

The following method depending on the hydrolysis of starch by hydrochloric acid and the subsequent estimation of the glucose produced, is only reliable if there are no pentosanes or other substances present which on hydrolysis would yield reducing sugars.

About 3 grams of the substance in as fine a state of division as possible are covered with 50 c.c. of cold water and shaken at frequent intervals; after an hour the insoluble portion is filtered off and washed with water until the total filtrate measures 250 c.c.; the addition of a little alumina shaken up with water will frequently facilitate clear filtration. The soluble carbohydrates contained in the filtrate may if desired be determined both before and after inversion.

The residue remaining on the filter paper is then transferred to a flask with a 250 c.c. graduation mark and heated for two and a half hours under a reflux condenser with 200 c.c. of water and 20 c.c. of hydrochloric acid (sp. gr. 1·125). After cooling,

^{*} Dennstedt and Voigtländer: "Ber. deut. chem. Gesells.," 1895, 28, R., 1025.

the solution is neutralized with caustic soda and made up to 250 c.c., whereupon it is filtered and the amount of glucose contained in an aliquot portion of the filtrate is estimated by Fehling's or Benedict's solution. The amount of glucose found when multiplied by org gives the weight of starch.

The following method for the estimation of starch in barley is due to Horace T. Brown *:—

Five grams of the powdered or crushed grain are extracted for three hours in a Soxhlet extractor with alcohol (sp. gr. 0'90); the residue is then thoroughly boiled with 100 c.c. of water, and, after cooling, to 57°, 10 c.c. of active malt extract are added and the mixture is set aside for one hour; it is thereupon boiled and filtered into a flask with a 200 c.c. graduation mark; the residue is thoroughly washed with water, and, after cooling, the filtrate and washings are made up to 200 c.c. The cupric reduction of 20 c.c. of the solution is determined under the conditions laid down by Brown, Morris and Millar,† the maltose being calculated according to Table XI in that paper (loc. cit., p. 100), after correction for the reduction due to the malt extract. The starch equivalent to this maltose is then ascertained by assuming that 84'4 parts of maltose correspond to 100 parts of starch.

The malt extract is prepared by digesting 10 grams of fresh finely-ground malt for two to three hours with 200 c.c. of water and filtering.

DEXTRINS.

The term dextrin is applied to substances which are polymeric with starch and are formed from it by the action of heat alone or of diastase or mineral acids. In the plant dextrins may occur as transitory substances whenever starch is being acted upon by diastase; further, certain dextrins may occur in a more permanent form. Thus the sap of the epidermal cells of *Arum italicum* turn reddish-violet on the application of iodine. The aqueous extract of such cells gives on evaporation a transparent sticky substance. This also gives with iodine a violet coloration; after boiling, the colour reaction

^{*} Horace T. Brown: "Trans. Guiness Research Lab.," 1903, 1, 89. † Brown, Morris and Millar: "J. Chem. Soc., Lond.," 1897, 71, 94.

with iodine is red, and after digestion with diastase a reducing sugar is found. A similar substance—termed soluble starch—has been described as occurring in the epidermis of Saponaria officinalis and also in Fungi. It must, however, be borne in mind that the glucoside saponarin,* $C_{21}H_{24}O_{12}$, is not uncommon in the epidermis of leaves of many plants, e.g. Saponaria officinalis itself, and as it gives a blue to violet coloration with iodine it is not unlikely that, in some cases, what has been described as soluble starch is really saponarin.

As already mentioned, the question of the formation of dextrins from starch by the action of diastase has been the subject of a great many researches, and has, at different times, resulted in the postulation of the existence of a large variety of dextrins and intermediate products, such as amylo-, achroo-, erythro-, and malto-dextrin, amylases, amyloins, glyco-amylins, etc., many of which did not survive for long.

The chief facts observed during the action of malt extract on starch may be very briefly summarized as follows. If, say, a 10 per cent starch paste is left in contact with malt extract at 50°, the mass rapidly liquefies and the solution acquires a sweet taste owing to the conversion of starch into maltose; if the latter substance be estimated from time to time, it will be found that the reducing power of the mixture increases rapidly at first until, after about two hours, the amount of maltose present corresponds to about 80 per cent of the starch employed, when practically no further change takes place. The change in the starch paste can also be demonstrated by periodic testing with iodine solution; the blue-black coloration gradually becomes less and less marked until various shades of red are obtained, finally the iodine gives no distinctive coloration. A corresponding fall in the optical activity of the solution can also be observed, but as the activity is still greater than what it should be for maltose alone, it must be concluded that some other substance is formed at the same time as the maltose, and that its reducing power is less but its activity is greater than that of maltose. The amount of this "non-maltose" product of diastatic activity varies directly with the temperature, and increases considerably at the expense of the maltose if the temperature be kept at or above 60°; if to such a pro-

^{*} Barger: "Ber. deut. chem. Ges.," 1902, 35, 1296.

duct, rich in non-maltose, a fresh quantity of malt extract be added, the non-maltose will be attacked and converted into maltose until the amount present again attains the value 80 per cent, which is the normal maximum; this experiment, which is due to Brown and Morris,* shows that the non-maltose is composed of different constituents, some of which are converted into maltose by diastase more readily than others; moreover experiments have shown that these substances behave differently towards yeast, some being more readily fermentable than others. This non-maltose constituent represents a mixture of the various dextrins mentioned above as having been described by several authors. More recently Maquenne and Roux † and others, carrying on the experiments of Brown and his collaborators, have found that on prolonged action extending over several days, even this non-maltose is slowly attacked by diastase, and a practically theoretical yield, instead of only an 80 per cent yield, can be obtained.

The action of malt on starch accordingly takes place in two stages, of which the first is rapid, being completed in about two hours, while the second one is very slow. According to Maquenne and Roux, the first stage corresponds to the hydrolysis of the amylo-cellulose (amylose) and the solution of the amylo-pectin with consequent formation of dextrins; the second or slow stage consists in the hydrolysis of these dextrins into maltose, and they consequently regard amylo-pectin as a maltosane.

It was mentioned above that a larger yield of non-maltose is obtained at higher temperatures, and that this is regarded as a mixture of dextrins, since some of it is readily converted into sugar on adding more diastase, whilst some still remains which resists; this latter is most likely produced from the amylopectin and corresponds to the stable dextrin described by Brown and Morris, whereas the easily converted portion is in all probability identical with what Brown and Morris called maltodextrin or amyloin, and may have been produced by a peculiar action of malt on the amylo-cellulose (amylose) constituent of the starch.

^{*} Brown and Morris: "J. Chem. Soc., Lond.," 1885, 47, 527. † Maquenne and Roux: "Compt. rend.," 1906, 142, 124, 1059.

General Properties of Dextrins.

From what has been said above, it will be seen that the term dextrin comprises a number of substances some of which are not at all well defined. The following may, however, be regarded as approximately representing the characteristics of all substances included in this group:—

- 1. They are amorphous substances which are readily soluble in water to form gummy solutions, which are used as a substitute for natural gum; they are precipitated from aqueous solutions by the addition of alcohol.
- 2. Unlike starch inulin and glycogen, dextrin does not give a precipitate with basic lead acetate.
- 3. As their name implies, they are strongly dextro-rotatory, in which respect they resemble starch.
- 4. They give either a red colour or no colour at all with iodine.
- 5. They are not fermentable by yeast alone, but are fermented by a mixture of yeast and diastase acting together, which is no doubt due to their slow hydrolysis in the first place by the diastase and the subsequent fermentation of the maltose so produced.
 - 6. They do not reduce Fehling's solution when pure.
- 7. They are converted into glucose on hydrolysis with mineral acids.

As has already been mentioned, starch when suddenly heated to about 200° is converted into a substance commercially known as dextrin. The use of starch for stiffening linen depends on some such similar change produced in the starch by the heat of the iron.

Although a great many different dextrins have from time to time been described, comparatively few of them are sufficiently well defined to warrant any description here. The three following, in addition to maltose and isomaltose, were isolated by Lintner and Düll* from the products of the action of malt extract on starch by a long process of fractional precipitation with alcohol.

Amylo-dextrin.—This substance, which is regarded by these authors as the chief constituent of soluble starch, is a white

^{*} Lintner and Düll: "Ber. deut. chem. Gesells.," 1893, 26, 2533.

powder which is slightly soluble in cold water, but readily in hot. It is strongly dextro-rotatory ($a_p = + 196$), does not reduce Fehling's solution, and gives a blue colour with iodine.

Erythro-dextrin.—This is a solid which dissolves readily in water, has a rotatory power of $a_{\rm b} = + 196^{\circ}$, and with iodine produces a red-brown colour.

The existence of erythro-dextrin as a chemical entity is, however, disputed by Ost, who says that it is a mixture of achroo-dextrin with starch; an artificial mixture of achroo-dextrin with a half per cent of starch also produces a red colour with indine

Activo-dextrin.—This substance is optically active, has the value $a_{\rm p}=+192^{\circ}$, gives no colour with iodine, and has a sweetish taste

COMMERCIAL DEXTRIN.

Commercial dextrin is prepared by heating starch to about 230-260°; it is a yellowish-brown powder, while that prepared by acid hydrolysis of starch is an almost colourless solid with a choncoidal fracture, or else a white powder resembling starch. It is composed chiefly of achroo-dextrin mixed with varying quantities of erythro-dextrin and glucose. It dissolves in an equal volume of water to give a neutral sticky solution with a faint sweet taste; the solution is strongly dextrorotatory. Dextrin is insoluble in alcohol and ether.

GLYCOGEN.

This substance, although one of the most important and widely distributed reserve foods in the animal kingdom, has a restricted distribution in plants. It occurs abundantly in certain Fungi, especially in *Saccharomyces cereviseae*, where it may sometimes form as much as 30 per cent of the dry weight. It has also been described as forming part of the cell-contents in Myxomycetes, Flagellates, and possibly in certain Algae and Cyanophyceae. In the yeast plant the glycogen varies in amount according to the physiological phase of the organism, and, it appears, accumulates and disappears often with great rapidity.

The glycogen appears in the cells of Saccharomyces during the early stages of fermentation as minute refractive granules scattered through the protoplasm; after a few hours these granules give place to small vacuoles, which in turn are replaced by one large vacuole, which may occupy the greater space in the cell. According to Harden and Rowland,* this progressive increase in the size of the glycogen-vacuole may result from the formation of some substance, besides carbon dioxide, from the glycogen.

Wager and Peniston† have shown that the amount of glycogen present is correlated with the periodical fluctuations in the fermentative activity.

On adding yeast to the nutrient fluid, e.g., Pasteur's solution, fermentation may start at once, in which case it was found that the cells float and contain very little glycogen. On the other hand, the cells may contain much glycogen and sink to the bottom; in this case fermentation is slow to commence, but gradually increases, and eventually becomes much more active; also the budding is much more extensive as compared with a yeast which contains but little or no glycogen.

If healthy brewers' yeast be added to Pasteur's solution the cells, which contain much glycogen, sink to the bottom. After an hour or two the cells begin to rise, and they become distributed throughout the medium after the lapse of four or five hours. The fermentation is now much more active, and the amount of glycogen in the cells less. The next five to fifteen hours is the period of maximum vegetative activity, during which the glycogen disappears; then it slowly reappears, and later on much more rapidly, at which phase there is a marked decrease in budding. At the height of fermentation, or immediately after, the glycogen increases rapidly, and a large number of cells sink to the bottom of the fluid. If the medium be not exhausted, the process may be repeated two or three times.

Although glycogen may be looked upon as a temporary reserve food-matter, for yeast-cells rich in glycogen retain their vitality much longer than those in which there is little

^{*} Harden and Rowland: "J. Chem. Soc., Lond.," 1901, 79, 1234. † Wager and Peniston: "Ann. Bot.," 1910, 24, 45.

or none, the fact that in the spores of species of *Mucor* and in sclerotia glycogen does not appear until growth has commenced, points to the conclusion that in these plants, at any rate, it is not primarily a storage product. Kohl considers that since it is more abundant in *Saccharomyces* during active gemmation, it is not exclusively a reserve substance, but an intermediate product in the formation of alcohol from the sugar.

In the animal kingdom, according to Hoppe-Seyler, glycogen is an invariable constituent of almost all developing cells; it is found also in the muscles and blood, and chiefly in the liver, where it is stored in larger quantities.

It may be remarked that there is little doubt that the glycogen obtained from animal and plant sources are identical.

Preparation.

The following method of obtaining glycogen was devised by Pflüger.* Fresh finely-cut liver is stirred up with water and 60 per cent caustic potash, and heated for two hours; the filtered solution, containing 15 per cent of potash, is then mixed with an equal volume of 96 per cent alcohol, and the precipitated glycogen is collected and washed with a mixture of one part of 15 per cent potash with two parts of 96 per cent alcohol; if necessary, the substance may be redissolved and purified in the same way.

Glycogen may also be prepared from yeast, but not in a particularly pure state, in the following manner: A quantity of bakers' yeast, which has been previously well washed with water, is mixed with fine well-cleaned sand and ground very thoroughly in order to rupture the cells. The mixture is then placed in a vessel with about thrice its volume of water and heated for some time, being constantly stirred. The liquid is then filtered off, cooled, and strong alcohol added to the filtrate in order to precipitate the glycogen, which is filtered off. The glycogen so obtained may be purified by redissolving it in water, adding a little acetic acid, and boiling in order to remove any proteins which may not have been removed by the initial heating, filtering, and precipitating with alcohol.

^{*} Pflüger: "Pflüger's Archiv f. Phys.," 1902, 91, 119, and 1903, 93, 163.

The following method has recently been described by Harden and Young*: The yeast is ground with an equal weight of sand. It is then extracted by boiling with water, and an equal volume of alcohol added to the cooled and filtered liquid. The precipitate formed is collected, washed with 50 per cent alcohol, and is then treated on the boiling water bath with a 60 per cent solution of potassium hydroxide for two hours. The liquid is cooled and poured into an equal volume of water, filtered, and the filtrate precipitated by the addition of two volumes of alcohol. The precipitate is collected, and washed repeatedly with a mixture containing 400 c.c. of water, 100 c.c. of 50 per cent potassium hydroxide, and 500 c.c. of alcohol; it is finally washed with 50 per cent alcohol.

The precipitate is then dissolved in water, and the solution, which is alkaline owing to the difficulty of washing away all the potassium hydroxide, is neutralized with acetic acid, and the glycogen precipitated by the addition of an equal volume of alcohol. By repeatedly dissolving in water, and reprecipitating with alcohol, a preparation may be obtained free from nitrogen and ash, but it still contains yeast-gum. which may be removed by redissolving the crude glycogen in water and saturating with ammonium sulphate. The precipitated glycogen, after being washed with saturated ammonium sulphate, is dissolved in water, and the solution again saturated with ammonium sulphate, the process being repeated three times. The final precipitate is again dissolved, the solution dialysed until free from ammonium sulphate, and the glycogen precipitated with alcohol. For details of the further purification of the glycogen the original paper should be consulted.

Properties.

Pure glycogen is a snow-white amorphous solid. It is readily soluble in hot water, forming an opalescent solution, from which it may be precipitated again by alcohol, provided small quantities of dissolved salts are present; 100 c.c. of a 1 per cent solution when mixed with 200 c.c. of absolute

^{*} Harden and Young: "J. Chem. Soc.." 1912. 101. 1928.

alcohol remain clear, but on adding 0.03–0.05 gram of sodium chloride, an immediate precipitate is formed. Glycogen is strongly dextro-rotatory, $a_{\rm p} = + 189.9^{\circ}$, and is coloured red to brown by iodine; it does not reduce Fehling's solution, but is broken up by diastase into dextrin and maltose, and by acids into glucose.

Identification.

- 1. The opalescent appearance of its aqueous solution is characteristic, and is strongly dextro-rotatory.
- 2. A brown coloration is given with iodine solution (cf. inulin, p. 119).
 - 3. A white precipitate is given with basic lead acetate.
 - 4. It does not reduce Fehling's solution.
- 5. On boiling with mineral acids, it is converted into dextrose.

Estimation.

This is best effected by heating the aqueous solution for three hours in a boiling water bath with about 2.2 per cent HCl, and then neutralizing and estimating the resulting glucose by means of Fehling's solution; the amount multiplied by 0.9 gives the weight of glycogen.

PARA-DEXTRANE AND PARA-ISODEXTRANE.

These substances have been isolated from *Boletus edulis* and *Polyporus betulinus* respectively. The former gives a yellow colour with chlorzinc iodide, and the latter a blue when treated with iodine and sulphuric acid. Both give glucose on hydrolysis.

LEVULOSANES.

INULIN.

This substance is commonly found as a reserve food-stuff, of the same nature as starch, existing in a state of solution in the cellsap of a number of plants belonging to the natural order Compositæ, e.g. in the tubers of the dahlia and artichoke (Helianthus tuberosus), and in the fleshy roots of the chicory (Cichorium Intibus). It has also been described as occurring in

the following natural orders: Violaceæ, Malpighiaceæ, Droseraceæ, Candolleaceæ, Goodeniaceæ, Campanulaceæ, Lobeliaceæ, Myoporineæ, Liliaceæ, and Amaryllidaceæ; also in some Algæ, e.g. *Neomeris*.

Inulin, or closely allied substances, are not infrequently found in company with starch, especially in some Monocotyledons; and the same peculiarity in its occurrence, as has already been remarked upon in connexion with the occurrence of starch in monocotyledonous plants, obtains (p. 97).

Thus in *Iris pseudacorus* starch is present but not abundant, in *Iris Xiphium* both starch and inulin are present in quantity; *Scilla nutans* has inulin but no starch, while *Scilla sibirica*, and also *Hyacinthus* and *Muscari botryoides* have both starch and inulin.

It is of interest to find that the nature of the reserve carbohydrates may often be correlated to the habitat of the plant. Parkin * points out that these reserve substances of aquatic plants and of plants inhabiting wet situations take the form of starch, e.g. Sparganium, Alisma, Listera, Orchis, and Schizostylis; whereas, on the other hand, inulin, generally associated with sugar, is the characteristic carbohydrate reserve in those Monocotyledons inhabiting dry situations, e.g. Allium, Asphodelus, Anthericum, Yucca, Tritona, Iris Xiphium, etc.

In this connexion \dagger reference must be made to the work of Lidforss, who showed that plants inhabiting wet situations fall into two distinct categories; those like *Elodea*, *Chara* and *Stratiotes*, which hibernate at the bottom of the pond or stream, contain starch but no sugar; while those which live on the banks where their rhizomes, or other organs of storage, pass the winter out of the water, e.g. *Myosotis* and *Menyanthes*, contain sugar during the winter months. In the former case a temperature of -2° C. to -4° C. is fatal, while in the latter case the death point is about -7° C.

This peculiarity also obtains for many arctic plants; Miyake, Wulff, and others have shown that cold, which means physiological dryness, is conducive to sugar production, so that arctic plants frequently exhibit but a small amount of starch, and relatively large quantities of sugar. Stahl has shown that

^{*} Parkin: "Phil. Trans. Roy. Soc., Lond.," B., 1899, 191, 169.

the leaves of mycotrophic plants, which generally show a feeble transpiration, seldom contain starch, its place being taken by glucose. Lidforss also has shown that the winter green vegetation of Sweden is characterized by the absence of starch from the leaves, the mesophyll, in its place, containing relatively large quantities of sugar, and sometimes oil during the winter months. In summer the leaves of these plants contain starch, which, on the advent of winter, is converted into sugar, from which starch is formed on the rise of temperature in the spring.

Then, again, it is not uncommon to find sugar stored in the periderm of trees and in the leaves of evergreen plants during the winter; starch, however, may be found in the leaves of evergreen trees during the cold season, its presence being due to feeble photosynthesis.

Reference may be made here to the well-known fact that potatoes turn sweet on exposure to cold. This conversion of starch into sugar is most active at 0° C., and the action decreases with the rise in temperature, so that above 7° C. no sugar is thus produced. Also if the tubers are suddenly subjected to a temperature of -1° C., no sugar will be produced. The amount of sugar formed is not great, its maximum being about 3 per cent of the wet weight; the limit of the process depends on the concentration of sugar, and, as Czapek has shown, the transformation of the starch may be prevented, on a lowering of the temperature, if the concentration of sugar be sufficient. If these sweet potatoes be exposed to a higher temperature, all the sugar that remains—some of course has been used up in respiration—is reconverted into starch.

Œcologically these characters are of value to the plant; for if the water of the cellsap be frozen, the salts held in solution become concentrated and will eventually precipitate the soluble proteins.

Parkin points out that the presence of inulin in the cell sap of the parenchymatous tissues would retard the evaporation of water. It is a well-known fact that water in the presence of oil may be much over-cooled before ice-formation takes place, and the freezing point of water in which other substances, e.g. sugar, are dissolved is depressed, and thus the danger arising from the salting out of the proteins is mini-

mized. But, notwithstanding these facts, plants are frequently subjected to temperatures sufficiently low to cause ice to be formed, and as the water is thus withdrawn, the sugar becomes more concentrated until it will also crystallize out. Both these processes generate heat, which may be sufficient in amount to enable the protoplasm to live. And this is, according to Mez and Lidforss, the explanation of the presence of sugar in winter leaves.

At the same time we must be careful not to push such explanations too far, for there are many exceptional cases; thus Ewart has pointed out that *Dicranum* which contains much oil is less resistant than is *Bryum*, and other mosses, in which such substances are absent. The beetroot also is very susceptible to cold, notwithstanding the fact that it contains much sugar; similarly the seeds of the hemp and willow, which contain much oil, are easily killed by desiccation, whereas the oil-containing seeds of the linseed are highly resistant. Such divergent phenomena must depend on the constitution of the protoplasm.

Again, oil is a convenient form of reserve food, especially in small organisms and in reproductive bodies, where space is limited and lightness is all-important and it is desirable to store a maximum of potential energy in the minimum of bulk. Finally, as Parkin points out, the nature of the carbohydrate reserve may depend on the kind of sugar transformed; thus, if saccharose be the chief sugar translocated from the leaves, then it might be expected that starch would be produced, on the other hand inulin might be formed if the available sugar for conversion were levulose.

Preparation.

Inulin may be obtained from dahlia tubers, of which it forms from 10-12 per cent, by crushing them and pressing out the liquid; the residue is then boiled up with a little water and some precipitated chalk and filtered again. The two filtrates are then united and once more boiled with chalk in order to neutralize any acids, and while still warm treated with lead acetate until no further precipitate is formed. The filtered solution is then saturated with hydrogen sulphide, fil-

tered, neutralized with ammonia, evaporated to half its bulk and mixed with an equal volume of alcohol. After one or two days, crude inulin may be filtered off; it may be further purified by warming in aqueous solution with animal charcoal, filtering and adding alcohol; the precipitated inulin is then washed with alcohol and ether, and dried over sulphuric acid.

According to Kiliani,* it may also be prepared by boiling crushed dahlia tubers with water and a little chalk, filtering and freezing the filtrate. When the water cools, the precipitate is filtered off, re-dissolved in hot water and frozen out once more. After repeating this process several times, the inulin is washed with methyl alcohol, ethyl alcohol, and finally ether.

Characters.

Pure inulin forms a white starchy tasteless powder of a sphæro-crystalline nature; it swells up and is readily dissolved in hot water, alkalis, etc., and may be recovered from the aqueous solution by the addition of alcohol, in which it is practically insoluble, or by freezing. Inulin is lævo-rotatory and unlike starch does not give a paste with water, nor does it give a blue colour when treated with iodine. Diastase has no effect upon it; it may, however, be hydrolysed by the ferment inulase, or by mineral acids, by which reagents it is converted into levulose. The low osmotic pressure which solutions of inulin exert suggests a large molecule, but its molecular structure appears to be less complex than that of starch. The relation between inulin and levulose is much the same as that existing between starch and glucose.

Identification.

In many plants the presence of inulin is indicated by the well-known sphæro-crystals which are obtained on steeping the fresh tissues for some time in strong alcohol; this deposition is not, however, always so characteristic; thus in Monocotyledons the inulin is frequently found, after treatment with alcohol, in amorphous masses. The sphæro-crystals and the amorphous concretions of inulin are readily soluble in warm

^{*} Kiliani: "Annalen," 1880, 205, 147.

water, and thus may be distinguished from calcium phosphate which may occur in cells in shapes similar to those of inulin. These two substances may be further recognized by the fact that sulphuric acid completely dissolves inulin, whereas it forms with calcium phosphate insoluble calcium sulphate. The following tests also may be performed,

- I. Green's Test.—Sections of the material, which have been soaked for some time in absolute alcohol, are treated with a saturated solution of orcin in strong alcohol, and then boiled in hydrochloric acid. The masses of inulin disappear and a red colour results. If phloroglucin be substituted for the orcin, the resulting coloration will be reddish-brown.
- 2. Molisch's Test.—The sections are treated with a 10 per cent alcoholic solution of a naphthol, then a few drops of strong sulphuric acid are added and the preparation warmed. A deep violet coloration ensues, and the inulin is dissolved.

These colour reactions are indicative of the formation of sugar by the hydrolysis of the inulin by the acids employed in the tests; it is therefore important, before employing these reactions, to make sure that no free sugars are present in the material to be examined, and to wash the preparations thoroughly with alcohol in order to remove them.

Since inulin does not reduce Fehling's solution, this reagent may be employed to ascertain whether any reducing sugars are present in the material before employing the above tests for inulin.

The following reactions may be carried out with a solution of inulin.

3. The addition of iodine solution gives with inulin a brownish coloration. Since the solution of iodine is itself brown, this test must be performed very critically. The following method may be employed: dilute the solution of iodine with water until it is a light brown colour, fill two test tubes with this solution and add to one a drop of the inulin solution; now compare the colour of the contents of the two test tubes.

This same reaction is also given by glycogen, when the same procedure may be followed.

4. Basic lead acetate gives with inulin, and also with glycogen, a white precipitate. This test may be used to distin-

guish inulin and glycogen from dextrin, which does not give a precipitate with this reagent.

- 5. Inulin is precipitated from solution by alcohol.
- 6. Hydrolyse with mineral acid and test for levulose.

There is as yet no very accurate method for the estimation of inulin. Dragendorff* recommends precipitating the inulin from an aqueous extract and then determining the amount of levulose which is produced on hydrolysis.

INULIN-LIKE SUBSTANCES.

Attention may now be drawn to substances similar to inulin which occur in various plants. The chief of these are:—

Graminin in Agrostis, Festuca, Trisetum and other grasses. Irisin in Iris pseudacorus.

Phlein in *Phleum pratense* and *Phalaris arundinacea*, Sinistrin in *Scilla maritima*.

Triticin in Triticum repens, Dracæna australis and Dracæna rubra.

All these compounds have the same formula, $6(C_eH_{10}O_5 + H_2O)$, and possess the same characteristics; they are lævorotatory, yield fructose on hydrolysis, and are fairly soluble in cold water. The majority are difficult to crystallize, and their solutions yield a gum-like substance on evaporation. It is possible that some, at any rate, of these substances may bear the same relation to inulin as dextrin does to starch.

MANNOSANES.

MANNANE.

The seeds of many plants contain reserve carbohydrate which is generally referred to as reserve cellulose, hemi-cellulose, or para-galactane substances. These materials are often indistinguishable from true cellulose by the microchemical means at our disposal. They may, however, be distinguished by the products of their hydrolysis; thus they form glucose, together with other substances, on treatment with hot dilute hydrochloric or sulphuric acid, whereas ordinary cellulose does not. Also they are dissolved by dilute alkalis, and by cuprammonia after

^{*} Dragendorff: "Jahres. Fortsch. d. Chem.," 1872, 929.

a *brief* treatment with hot dilute hydrochloric acid. These reserve celluloses contain mannanes and galactanes.

Mannane serves in the same way as starch as a reserve foodsupply to a very large number of different plants. It may be regarded as an anhydride of mannose, since it yields this substance on hydrolysis; it occurs either alone or united to anhydrides of other sugars, such as glucose, galactose, pentose, etc., in a great many different forms.

A fairly pure specimen of mannane can be obtained from yeast by a somewhat elaborate method devised by Hessenland.* It is a white amorphous substance, which is somewhat soluble in water and swells up in dissolving; it is insoluble in alcohol but readily soluble in alkali, and is strongly dextro-rotatory $a_p = +283.7-287.6$.°

Mannane occurs in salep mucilage, and has been extracted by Ritthausen† and Effront‡ and others from wheat and barley. Mannanes are also found in *Penicillium glaucum*, ergot, in the roots of several plants such as asparagus, chicory, *Helianthus* and *Taraxacum*; also in the wood and leaves of many trees, such as lime, chestnut, apple, mulberry, certain Oleaceæ and conifers; the so-called reserve celluloses and hemi-celluloses contained in seeds of Palmaceæ, Liliaceæ, elder, cedar and larch, and many other plants, are also very rich in mannanes.

PARAMANNANE.

Paramannane is a variety of mannane which is characterized by being much more resistant to hydrolysis; this substance, which is contained in coffee beans, is only slightly acted on by hot dilute mineral acids, potassium chlorate and hydrochloric acid, but dissolves in a concentrated hydrochloric acid solution of zinc chloride. It is accordingly frequently classed as a mannose-cellulose.

CARUBIN OR SECALANE.

Carubin § is the name given to a substance occurring in the seeds of *Ceratonia siliqua*, and in various cereals such as rye

^{*}Hessenland: "Z. d. Vereins d. Deut. Zuckerind.," 1892, 42, 671.

⁺ Ritthausen: "J. prakt. Chem.," 1867, 102, 321, and "Chem. Zeit.," 1897

[‡]Effront: "Compt. rend.," 1897, 125, 38, 116.

[§] Ibid., 124, 200, and 125, 116 and 309.

and barley. In its characters it closely resembles mannane, and by some authors is regarded as identical with it; when dry, it is a spongy friable substance which swells upon the addition of water. It is soluble in cold water and is optically inactive. Its sugar is fermentable and non-crystalline.

GALACTOSANES.

GALACTANE.

Exactly analogous to the mannanes are the galactanes, which may be looked upon as anhydrides of galactose. They occur in a great variety of different forms, some of which are readily hydrolysed by warming with alkali, while others are very resistant even towards boiling alkali. Four galactanes have been described, which are distinguished by the prefixes α -, β -, γ - and δ -; they are all amorphous substances which dissolve with difficulty in water, and on hydrolysis yield galactose.

PARAGALACTANE.

Paragalactane is a substance which is better termed paragalacto-arabane, since on hydrolysis by weak mineral acids it yields a mixture of galactose and arabinose. It occurs in the cell walls of the cotyledons of many plants, e.g., *Lupinus luteus* and other species, *Phænix dactylifera*, *Cocos nucifera*, and other palms, *Soja hispanica* and *Coffea arabica*, where it forms a reserve food-material which is digested on germination.

Paragalactane is a white solid which is insoluble in water and cuprammonia; it dissolves in hot potash. On heating with nitric acid it is oxidized to mucic acid. Microchemically it may be identified by its insolubility in the reagents mentioned, and also by the fact that with phloroglucin and hydrochloric acid it gives a red coloration on warming. No colour is given in the cold.

Its association with cellulose prevents the latter exhibiting some of its reactions; thus the cellulose is unacted upon by cuprammonia unless the paragalactane be removed; this may be done by boiling in dilute hydrochloric acid.

GUMS 123

AMYLOID.

Amyloid* is the name given to a substance occurring in the seeds of pæonies and certain cresses,† which yields on hydrolysis with dilute sulphuric acid a mixture of galactose, glucose, and xylose. It is a colourless substance, and is insoluble in cold water, but swells up into a slimy mass in hot water; it is soluble in cuprammonia solution. Amyloid does not reduce Fehling's solution, but is oxidized by nitric acid to mucic and trihydroxy-glutaric acids. It gives a blue colour with iodine.

GUMS.

The natural gums were formerly thought to be carbohydrates of the general formula $(C_eH_{10}O_{\delta})_n$; the researches of O'Sullivan, however, have shown that they are not simple carbohydrates, but are rather substances of a glucosidal nature, since on hydrolysis they give rise to sugars mixed with complex acids of high molecular weight. The gums themselves retain the character of acids, and it would appear that the molecule of a gum is composed of a number of sugar residues grouped around a nucleus acid in such a way as to leave the acid group exposed.

The gums are translucent amorphous substances, some of which dissolve in water completely, giving a sticky solution, while others merely swell up in water and form a sort of jelly; they are all insoluble in alcohol.

The natural gums must be distinguished from starch gum or dextrin, which is an artificial product obtained from starch, by the following characteristics:—

- I. Solutions of natural gums are lævo-rotatory, whereas those of dextrin are dextro-rotatory.
- 2. Basic lead acetate precipitates natural gums from solution, but has no action on dextrin.
- 3. Natural gums on hydrolysis yield chiefly galactose and pentoses such as arabinose or xylose, whereas dextrin yields glucose only.

The hydrolysis of gums takes a long time to complete-

^{*} Cf. footnote, p. 135. + Winterstein: "Z. physiol. Chem.," 1893, 17, 353.

from eighteen to twenty-four hours—whereas dextrin is easily hydrolysed.

4. On oxidation with nitric acid, natural gums yield chiefly mucic acid $(C_eH_{10}O_8)$ together with some saccharic $(C_eH_{10}O_8)$ and oxalic $(C_2H_{20}O_4)$ acids, whereas dextrin yields chiefly oxalic acid together with a small quantity of saccharic and tartaric $(C_4H_6O_6)$ acids.

As they occur in nature, the true gums are mostly combined with potassium, calcium, or magnesium in the form of salts, from which the true carbohydrate can be isolated by the action of a stronger acid.

The classification of gums is, for want of more accurate knowledge, based chiefly on their solubility in water:

- (a) Gums, such as arabin, which are completely soluble.
- (b) Gums which are partially soluble, such as cerasin and bassorin.
- (c) Mucilages and pectic bodies which merely swell up with water to form a jelly.

The classification, however, is by no means rigid, many natural gums being composed of mixtures of several kinds of gums.

In the separation of gums from the tissues of the plant, advantage is taken of their solubility in water; it is found in practice, however, that in many cases mere maceration in water does not remove all the gum present; Dragendorff found that much more arabic acid could be extracted after the material had been treated with an alcoholic solution of tartaric acid.

Microchemical Reactions.

Microchemically, gum and mucilage may be recognized by their solubility and swelling respectively in water. Both are insoluble in alcohol and ether. With other reagents the results differ in different examples. Thus with iodine either a blue or a yellow colour may result, while in other cases the blue coloration is only obtained after treatment with chlorzinc iodide or sulphuric acid and iodine. Then again different degrees of solubility are found to obtain on treatment with

cuprammonia. Many of these substances stain well with corallin soda, and they also, especially the mucilages, show a great avidity for stains such as aniline blue and aniline violet.

GUM ARABIC.

This substance is a mixture of calcium, magnesium, and potassium salts of a weak acid of unknown constitution, to which earlier writers gave the name of arabic acid or arabin. O'Sullivan, however, applied the term arabic acid to a substance of the formula C₂₂H₂₂O₂₂, which he regarded as the nucleus acid around which a number of sugar residues are grouped: by hydrolysis under varying conditions, it is possible to split off successive sugar residues with the formation of acids of gradually decreasing molecular weight, until finally the nucleus acid free from all carbohydrate residues remains, and it is this acid that he calls arabic acid: the natural gum itself would, according to him, be a diarabinan-tetragalactanarabic acid of the formula ${}_{2}C_{10}H_{10}O_{8}$, ${}_{4}C_{19}H_{20}O_{11}$, ${}_{23}H_{30}O_{18}$, which is combined with the calcium, magnesium, and potassium. The arabic acid of the earlier authors, which is the acid set free from the natural gum by the removal of the calcium, magnesium, and potassium, may be prepared by acidifying a concentrated aqueous solution of gum arabic with hydrochloric acid, and adding alcohol. The pure substance is a white amorphous glassy mass which dissolves in water to give a lævo-rotatory solution. Ten per cent sulphuric acid converts this arabic acid into metarabic acid, which swells up in water, but does not dissolve.

Reactions.

Solutions in water (10 per cent) of arabic acid and other varieties of gum arabic give, according to Masing,* certain more or less definite reactions.

- I. They are not precipitated by (a) a cold saturated solution of copper acetate; (b) 10 per cent solution of lead acetate; (c) solution of ferric chloride (sp. gr. 1.2).
 - 2. A five per cent solution of silicate of potash produces a

^{*} Masing: "Archiv d. Pharm.," 1879, [3], 15, 216; 1880, 17, 34, 41; "Year Book of Pharmacy," 1881, 191.

cloudiness or a precipitate which is partially or wholly soluble on adding an excess. Arabic acid either does not respond to this reagent, or merely gives a slight turbidity, and the same applies to the gums obtained from certain species of *Cactus*, *Albizzia*, *Acacia catechu*, *Acacia leucophlwa* and other plants.

- 3. Stannate of potash gives similar reactions, and in the case of arabic acid produces a precipitate which is soluble in excess.
- 4. A solution of neutral sulphate of aluminium (10 per cent) generally gives a precipitate which is, in many cases, soluble in potash.
- 5. Basic lead acetate yields a precipitate which is entirely or partially soluble in excess.

GUM TRAGACANTH.

This gum occurs in species of *Astragalus*, and consists of about 8-10 per cent of soluble calcium, magnesium, and potassium salts, together with about 60-70 per cent of insoluble salts, which only swell up in water to a jelly. The water soluble portion is said to contain a substance, polyarabinon-trigalactan-geddic acid, which on hydrolysis breaks up into arabinose, galactose, and geddic acid, an isomer of arabic acid. The part soluble in water, when treated with baryta water, gives two isomeric tragacanthan-xylan-bassoric acids, which on hydrolysis yield a pentose sugar tragacanthose, xylose, and bassoric acid $C_{14}H_{20}O_{13}$.

WOOD GUM AND CERASIN OR CHERRY GUM.

These are other examples of pentosanes, but too little is known of their chemistry to warrant any description.

WOUND GUM.

A gum-like substance, termed wound gum, is frequently found in the tracheæ of plants, in the immediate neighbourhood of wounds, and stopping up the lumina; it is secreted by the surrounding living cells. Wound gum does not swell in water, and is insoluble in sulphuric acid and in caustic soda. On oxidation with nitric acid it yields both mucic and oxalic acids, and it responds to lignin tests; e.g., on treatment with

phloroglucinol and hydrochloric acid a bright red coloration results.

MUCILAGE.

The term mucilage is applied to those substances which with water produce a slimy liquid. Mucilage is widely distributed, and occurs in all or nearly all classes of plants. Mucilage-secreting hairs, or comparable structures, occur in various Muscineae, Filices, and especially in the Phanerogams; mucilage sacs or canals are found in certain Muscineæ, e.g., Anthoceros, Marattiaceæ, some Cycadaceæ, and Phanerogams; finally, the external walls of plants may be generally mucilaginous; e.g., in very many Algæ, the hibernaculæ of some aquatic Phanerogams, like Utricularia and Myriophyllum, and finally in the coats of seeds and fruits, such as Lepidium and Sterculia scaphigera respectively, in which cases the superficial cell walls are mucilaginous. Mucilage is not infrequently associated with other substances; thus in the case of mucilagesecreting hairs, it is sometimes associated with tannin, and in many plants, especially in the mucilage sacs of many Monocotyledons, calcium oxalate is found.

The constitution of mucilages is as yet unknown; they are, however, related pretty closely both to cellulose and to arabin. In fact, by some authors they are regarded as decomposition products of cellulose, produced either by overnutrition of certain cells or by bacterial action; * according to Wiesner, all gums are produced by a diastatic ferment acting on cellulose; although it is not possible to express any definite views on the subject, it would appear not improbable that in many cases the formation of gums and gum-like substances in the plant is a morbid condition. Mohl was able to show in the case of tragacanth gum that this substance was produced by the metamorphosis of the cells of the medullary rays.

That mucilages are not all of the same constitution is shown by the fact that the mucilaginous substance obtained from *Fucus* (caragheen mucilage) on hydrolysis with dilute sulphuric acid yields galactose, while salep mucilage, obtained from *Orchis Morio*, on a similar treatment yields mannose.

^{*} See Greig Smith: "J. Soc. Chem. Ind.," 1904, 105, 972.

Function.

Mucilage, when it is a definitely secreted product or of a definite and constant occurrence in a plant, may perform several functions, but how far these are primary functions cannot yet be stated.

When it occurs in tubers, as in the Orchidaceæ, mucilage is generally looked upon as a reserve food-material; it may serve as a check against too rapid transpiration, especially when produced in connexion with developing organs, such as vegetative buds, young leaves, in the epidermis of mature leaves, the sporangia of Cryptogams, etc.; in the case of aquatic plants, such as Algæ, the hibernaculæ of Myrio-phyllum, etc., its presence may prevent a too rapid diffusion; the calcareous incrustation of certain Algæ, e.g., Neomeris dumetosa, is dependent on the presence of mucilage; mucilage provides a water-storage mechanism in plants subjected to xerophytic conditions, e.g., Cassia obovata, Malva parviflora, Theobroma cacao, and Pterocarpus saxatilis; finally, it may be an important aid in connexion with seed-dispersal and germination, as in some species of Salvia and Lepidium.

Related to the gums and mucilages are the substances known as galactosanes occurring in the seeds of Leguminosæ (*Lupinus*, *Medicago*, etc.); wood gum or xylane, occurring in wood, etc. etc. These substances have already been mentioned in connexion with the sugars which they give rise to on hydrolysis.

PECTIC BODIES.

Many succulent fruits, such as pears, apples, gooseberries, and currants, and also fleshy roots, such as carrots, beetroots, etc., contain, together with the cellulose in the cell walls of parenchymatous elements, a substance which is soluble in water, but whose aqueous solution gelatinizes on the addition of alcohol. This substance, which is also probably the cause of concentrated aqueous extracts of fruit gelatinizing, is known as pectin.

According to Frémy,* the hardness of unripe fruit is due

^{*} Frémy: " J. Pharm. et Chim.," 1840, 26, 368.

to the presence of a substance known as pectose, which is deposited in the cell walls; as the fruit ripens the pectose undergoes a variety of changes, and is ultimately converted into pectin.

Under the action of an enzyme pectase contained in the plant, pectin is coagulated; this change was first studied by Frémy, and later by Bourquelot and Hérissey; * according to Duclaux † and others, the coagulation is dependent on the presence of calcium salts, and will take place even in the absence of the enzyme.

Comparatively little is known about the chemistry of pectin or the pectic bodies, as there appear to be several of these substances; at one time there was even some doubt as to whether they were really carbohydrates, since the ratio of hydrogen to oxygen seemed to be less than that required for compounds belonging to this group. Analyses by Tromp de Haas and Tollens, however, agree fairly well either for the formula $(C_6H_{10}O_8)_n$ or $2C_8H_{10}O_5$. H_2O .

By boiling pectose with dilute acids or caustic alkalis, a number of different substances are produced, such as pectin, parapectin, metapectin, pectic acid—which is combined with bases, such as calcium, and forms the middle lamella of cell walls—parapectic acid and parapectosic acid, some of which are soluble in water, while others, such as pectin, swell up in water and gelatinize. The final product of these changes, namely, metapectic acid, is readily soluble in water; it would appear to be closely related to, or identical with, arabane, and on hydrolysis with dilute sulphuric acid gives arabinose.

This view receives confirmation from the work of Bourquelot and Hérissey, who have discovered an enzyme occurring in malt, which is not identical with diastase, and which is capable of hydrolysing pectose to a reducing sugar, namely, arabinose. This enzyme, to which they gave the name pectinase, acts both on unaltered and on coagulated pectic bodies, but, conversely, the coagulating enzyme pectase is without

^{*} Bourquelot and Hérissey: "J. Pharm. et Chim.," 1898, [6], 8, 145; 1899, [6], 9, 563, and 10, 5.

⁺ Duclaux: "Traité de Microbiologie," 1899, 11, 336, and Goyaud: "Comptrend.," 1902, 135, 537.

[‡] Tromp de Haas and Tollens: "Annalen," 1895, 286, 278.

action on pectic bodies which have been previously hydrolysed by pectinase.

Microchemical Reactions.

The fact that these pectic substances are akin to cellulose, and occur in conjunction with it, renders its identification by microchemical means somewhat difficult. Mangin * more particularly has investigated these matters, and gives the following methods:—

- I. Methylene blue, Bismarck brown, and fuchsin stain pectic substances, lignified and suberized walls, but not pure cellulose. If sections thus stained are treated with alcohol, glycerine, or dilute acids, the lignified or suberized walls retain their coloration, whilst the pectic substances are decolorized with rapidity.
- 2. Crocein and nigrosin stain lignified and suberized walls, but do not stain pectic compounds.
- Crocein, naphthol black, and orseille red stain pure cellulose, but do not stain pectic substances; similarly, pectic compounds are unstained by congo-red and azo-blue, whilst cellulose and callose are.
- 4. The middle lamella, which consists of compounds of pectic acid, may be differentiated from the other pectic substances which are mixed with the cellulose of the cell walls by the following method: A thin section is placed in a 20-25 per cent solution of hydrochloric acid in alcohol for twenty-four hours; the section is then washed with water and treated with methylene-blue or phenosafranin. The middle lamella stains much more deeply than the rest of the wall.
- 5. If, after the above treatment with acid alcohol, the section be washed in a 10 per cent solution of ammonia, it is found that the cells separate with ease one from the other. According to Mangin, the combined pectic acid is freed from its bases by the treatment with acid alcohol, and is then dissolved by the ammonia. A recombination of the pectic acid may be brought about by treatment with baryta water, and after this process the cells will not separate one from the other.
- 6. The cellulose may be separated in the following manner: A thin section is treated with cuprammonia for twenty-four

^{*} Mangin: "Compt. rend.," 1889, 109, 579; 1890, 110, 295, 644,

hours; it is then washed, first with water, and, finally, with 2 per cent solution of acetic acid. The cellulose is thus dissolved and fills the cells and intercellular spaces. On treatment with chlorzinc iodide the middle lamella gives either no colour reaction or turns a pale yellow, while the cellulose gives the familiar blue reaction; the membrane stains very deeply with safranin or methylene blue, and is easily soluble in a solution of ammonium oxalate.

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O'Su!livan: "J. Chem. Soc., Lond.," 1884, 45, 41; 1890, 57, 59; 1891, 59, 1029; 1901, 79, 1164.

H. Robinson: "Brit. Ass. Reports, York," 1906, 227.

Haynes: "Biochem. Journ.," 1914, 8, 553.

CELLULOSE.

The term cellulose should be taken in general to connote a group of substances rather than a single chemical compound; used in this generic sense, it comprises a number of substances of somewhat different origin and somewhat different characters. whose chief common properties are their physiological origin and their function in forming the basis of the material which is isolated by the protoplasm of the living cell for the purpose of forming the wall or periphery of that cell. Though met with chiefly in the vegetable kingdom, its occurrence in the animal kingdom is not unknown, since a substance described as Tunicin. said to be identical with cellulose, has been found in the cell walls of certain Tunicates and insects. In the course of time the cellulose originally formed is altered by the addition to it of various secondary products known as encrusting substances; thus the process of lignification consists in the conversion of cellulose into ligno-cellulose; accompanying this change is a gradual disappearance of the protoplasm. Thus the protoplasm within the cell produces a number of different substances which are deposited in the cell wall, the nature and properties of the resulting fibre depending, of course, on the nature of these substances.

CLASSIFICATION OF CELLULOSES.

The naturally occurring celluloses may be divided into the following groups:—

- I. Typical or Normal Celluloses of the Cotton Type.— These are exemplified by the cellulose obtained from cotton, flax, hemp, etc.
- II. Compound Celluloses of the Wood Cellulose, Jute and Cereal Grass Types.—The natural celluloses occurring in jute, cereal straws, esparto grass, etc., consist of some form of cellulose combined with a non-cellulose constituent, which may be either of the nature of lignin in the case of lignocelluloses, or a pectic or gummy substance in the case of pectocelluloses, or a fatty substance in the case of adipocelluloses. This group may therefore be subdivided into—
 - (a) Lignocelluloses.
 - (b) Pectocelluloses.
 - (c) Adipo- or Cuto-celluloses.

III. Hemi-, Pseudo- or Reserve Celluloses.—This is a somewhat heterogeneous collection of substances which differ structurally from the fibrous celluloses, and occur in the cell walls of the seeds of various plants; such as Coffea arabica, Soja hispida, Lupinus luteus, Cocos nucifera, Tropæolum majus, Impatiens balsamifera, Pæonia officinalis, and in peas and beans; celluloses of this type are much more easily hydrolysed than other celluloses, and give rise to various sugars, such as mannose, galactose and pentose. For this reason they may be regarded as anhydrides of these sugars, and are therefore treated under the heading of mannosanes (p. 120), galactosanes (p. 122), and pentosanes (pp. 57, 126).

In this group of celluloses are also included those which, according to the researches of Brown and Morris, are dissolved by the enzymes secreted by the germinating seed; these are sometimes referred to as reserve cellulose, though the name seems ill-chosen, inasmuch as they would not appear always to function as reserve material.

One of the richest sources of cellulose in nature is the cotton plant. The following table, taken from Bowman,* represents approximately the composition of cotton fibre from various sources.

^{*} Bowman: "The Structure of the Cotton Fibre," London, 1908, p. 147.

Source of Cotton.	Surat.	American.	Egyptian.
Cellulose	Per cent. 91.35 *40 *53	Per cent. 91.00 '35 '53	Per cent. 90.8 .42 .68
Mineral matter, i.e., salts of K, Na, Ca, Mg, Fe, and Al	*22	12	*25
Water	7.20	8.00	7*85

Such cellulose, however, in its native condition is not in a pure state, being in more or less intimate chemical union with other substances, such as pectic bodies, lignocellulose and colouring matters from which it has to be freed by a series of successive chemical treatments before the pure cellulose can be isolated

The chemical treatments referred to are as follows:--

- 1. Alkaline hydrolysis, which consists in boiling the fibres with 1-2 per cent caustic potash, and washing to remove the pectic bodies.
- 2. Exposure of the washed fibres to bromine or chlorine at the ordinary temperature; by this process the lignone complex of the lignocellulose is destroyed.
- 3. A second alkaline hydrolysis with sodium sulphite, carbonate or hydrate.

The cellulose is thus isolated in a very pure state.

CHARACTERISTICS AND PROPERTIES OF NORMAL CELLULOSE.

In describing the chemical properties of cellulose, the cellulose isolated as above described from the fibre substance of cotton is chosen as typical.

Pure cellulose is a white hygroscopic substance, which absorbs about 6-12 per cent of water, which it loses again when heated to 100° ; it is insoluble in water at ordinary pressure, but when heated with water in sealed vessels at 500° F., it is dissolved completely with decomposition.

SOLUBILITY OF CELLULOSE.

Cellulose is insoluble in all ordinary solvents, but when treated with zinc chloride in the presence of water, it is converted into a gelatinous hydrate which, after prolonged treatment, goes into solution.

A solution of six parts of zinc chloride in ten parts of water heated to 60-100° is thoroughly stirred up with one part of cellulose, and then digested for some time at a gentle heat. When the cellulose is gelatinized, its solution is completed by heating over a boiling water bath, and adding water from time to time to replace that lost by evaporation.

- Two other salt solutions are known which dissolve cellulose:—
- (a) Zinc chloride and hydrochloric acid.—A solution of zinc chloride in twice its weight of hydrochloric acid dissolves cellulose rapidly in the cold.
- (b) Ammoniacal cupric oxide (Schweitzer's Reagent).—The solution is prepared by adding ammonium chloride and then excess of sodium hydrate to a solution of a cupric salt; the blue precipitate so obtained is then washed, pressed on a cloth filter, and dissolved in 0.92 ammonia. Cellulose dissolves in this solvent and on the addition of acid is reprecipitated; this fact is made use of in the preparation of artificial silk.

ACTION OF VARIOUS CHEMICALS ON CELLULOSE.

I. Alkalis.—Solutions of caustic soda of I to 2 per cent strength have no action on cellulose at temperatures considerably above 100°. Solutions containing 10 per cent have a curious effect on cotton fibre, causing it to thicken and become more cylindrical, and destroying the central canal. This phenomenon was first made use of technically by Mercer, who found that by this means cotton could be made to acquire a gloss resembling that of silk, since the fibre becomes translucent during the contraction.

When fused at 200-300° with a mixture of sodium and potassium hydroxides, cellulose undergoes complete decomposition with the formation of oxalic and acetic acids.

The so-called alkali cellulose obtained by mercerizing cellulose with about 15 per cent caustic soda reacts with carbon disulphide to form xanthogenates;* these compounds are used in the manufacture of viscose (see below).

^{*} Cross, Bevan and Beadle: "Ber. deut. chem. Gesells.," 1893, 26, 1090; and Cross and Bevan: "Ber. deut. chem. Gesells.," 1901, 34, 1513.

2. Acids.—Nitric acid (sp. gr. 1.25) at 180° converts cellulose into axycellulose, a substance of a weak acidic character, which reduces Fehling's solution (see below under oxidizing agents). Concentrated nitric acid, or a mixture of this acid with concentrated sulphuric acid, converts cellulose into nitrates, the composition of which varies with the conditions of the experiment; di-, tri-, tetra-, penta- and hexa-nitrates,* which are of considerable technical importance, are known. Dilute sulphuric acid on prolonged action converts cellulose into hydro-cellulose, a substance of the formula $C_{12}H_{22}O_{11}$; this substance retains the structure of the cotton fibre from which it is produced, but on rubbing it breaks up into a fine powder. The same substance may also be obtained by the action of aluminium or magnesium chlorides at a temperature of 300° F.

Concentrated sulphuric acid dissolves cellulose, gradually converting it into dextrin and ultimately into dextrose. If the solution as soon as made is diluted with water, a gelatinous hydrate is precipitated; † this substance is known as amyloid, the since it resembles starch in giving a blue colour with iodine. The same substance is formed by the action of chlorzinc iodide, the reaction being used as a test for cellulose.

The combined action of glacial acetic acid and acetic anhydride in the presence of concentrated sulphuric acid or zinc chloride converts cellulose in *acetyl cellulose*, which is insoluble in water but soluble in several organic solvents. Acetyl cellulose is also used in the manufacture of artificial silk.

Cellobiose, $\S C_{12}H_{22}O_{11}$, is a disaccharide obtained in the form of its acetate by acting on cellulose with acetic anhydride and concentrated sulphuric acid. It stands in the same relation to cellulose as does maltose to starch; since cellulose and starch yield different disaccharides on hydrolysis, it would appear that these two substances are fundamentally different and that cellulose is not a higher polymer of starch.

^{*} See footnote, p. 140.

[†]This reaction is made use of in the preparation of parchment paper. For this purpose paper is rapidly drawn through a mixture of four parts of concentrated sulphuric acid with one part of water; the paper is then thoroughly washed with water until it is free from acid.

[‡]This substance must not be confused with a compound of the same name which occurs naturally in several plants (cf. p. 123).

[§] Skraup and König: "Ber. deut. chem. Gesells.," 1901, 34, 1115; Schliemann: "Annalen," 1911, 378, 366.

Cellobiose reduces Fehling's solution and gives an osazone, m.p. 208-210°.

3. Oxidizing Agents.—Dilute solutions of alkaline hypochlorites have very little action on typical cellulose, and can therefore be employed for bleaching this material; with concentrated solutions of hypochlorites, however, a general decomposition ensues. As already mentioned, nitric acid (sp. gr. 1.25) at 180° converts cellulose into a series of oxidation products known as oxycellulose, and similar substances are produced by the action of other oxidizing agents, such as chromic acid, potassium chlorate, and hydrochloric acid, etc. The nature of these oxycelluloses differs somewhat according to their mode of formation, but in general they are characterized by the fact that they yield a relatively large amount of furfurol on boiling with hydrochloric acid; they are hydrolized by boiling with milk of lime into isosaccharic and dioxybutyric acids; they also reduce Fehling's solution, and are dyed by basic dyes, such as methylene blue.

The fact that the cellulose obtained from esparto grass and cereal straws resembles oxycellulose, in yielding a considerable proportion of furfurol on boiling with hydrochloric acid, leads to the idea that cellulose from these sources contains oxycellulose, but whether or not such oxycelluloses are actually pre-existent in the plant fibre has not yet been definitely established (see Lignocelluloses).

4. Action of Ferments.—It has been shown by Brown and Morris, in the case of malt, that the cell wall of the endosperm cells which contain nutrient material are broken down by a cellulose-dissolving ferment, a cyto-hydrolyst, before the embryo can procure the food-stuff contained in these cells. This enzyme, which is developed during the germination of the seed, can be extracted from the malt by cold water, and precipitated from this solution by alcohol. As another example of the fermentative decomposition of cellulose may be quoted the formation of marsh gas according to the equation

$$C_6H_{10}O_5 + H_2O = 3CO_2 + 3CH_4$$

which may be observed when vegetable matter is undergoing slow decomposition under stagnant water.

CHARACTERS AND PROPERTIES OF COMPOUND CELLULOSES.

As already stated, the main characteristic of the group of compound celluloses is that they are composed of one or other form of cellulose combined with some other substances of a non-cellulose nature.

The nature of the *cellulose constituent* varies according to the source from which it is obtained, one of the chief characteristic differences between such different forms of cellulose being their behaviour on boiling with hydrochloric acid; thus whereas cotton cellulose yields only about 0·1-0·4 per cent of furfurol, jute cellulose under similar conditions yields 3·0-6·0 per cent, and straw cellulose yields from 12·0-15·0 per cent; for this reason the cellulose constituent is regarded as being of the nature of oxycellulose.

The Non-cellulose Constituent of compound celluloses may vary very considerably in chemical nature, and on this fact depends their classification into—

- (a) Lignocelluloses.
- (b) Pectocelluloses.
- (c) Adipo- or Cuto-celluloses.
- (a) Lignocelluloses.—In the young cell the walls consist of almost pure cellulose, but, as the cell grows older, the walls may become permeated with what are known as encrusting substances, the process being known as lignification. This change takes place at the expense of the cellulose, and a new substance known as lignocellulose is produced. The extreme limit of this change is the production of wood, which contains only about 50-60 per cent of cellulose, while lignocelluloses still contain about 70-80 per cent.

These lignocelluloses are considered by most authors to consist of cellulose combined with at least two other non-cellulose constituents; one of these, A, appears to contain an aromatic nucleus, and the other, B, contains a furfurol-yielding complex, and is probably a pentosane. The two constituents, A and B, are sometimes grouped together as a single substance under the name of lignin or lignone. The constitution of this substance is still unknown. According to Klason* lignin would appear to be related to coniferyl alcohol (p. 189).

^{*} Klason: "Arkiv. Kem. Min. Geol.," 1908, 3, No. 5, 1.

It may here be remarked that although the best quality paper is manufactured from cellulose freed as completely as possible from non-cellulose constituents by the method described below, lignocelluloses are used as such for the preparation of inferior qualities of paper, without previous treatment for the removal of the lignin. Such papers when dipped in a 1½ per cent alcohol solution of phloroglucinol and touched with a drop of diluted hydrochloric acid are coloured red. They are, moreover, turned yellow by exposure to sunlight.

- (b) Pectocel'uloses, which occur in fibrous and parenchymatous tissues of the stems and roots of Phanerogams, are substances in which the non-cellulose constituent is of a colloidal or gummy nature and belongs to the group of substances known as "Pectic Compounds" (see p. 128). Similarly the mucocelluloses which are found in the seeds and fruits of Phanerogams and in Algæ contain for their non-cellulose constituent substances of a mucilaginous or gummy nature. The chemical characteristics of the pecto- and muco-celluloses are not, however, sufficiently well defined, as yet, to warrant a detailed consideration.
- (c) Adipo- and Cuto-celluloses.—Information regarding the composition of the constituents of suberized and cuticularized walls is very meagre. Till quite recently these compounds have been looked upon as cellulose in association with substances of a fatty or wax-like nature, and known as suberin and cutin. This view is based upon the fact that suberized walls, if treated first with a solution of potash, turn with chlorzinc iodide a red-violet colour.

The work of Gilson* tends to show that cellulose does not enter into the composition of such walls for the following reasons:—

- 1. Cellulose is not attacked by prolonged boiling in a 3 per cent solution of potassium hydrate in alcohol; suberized walls, on the other hand, are dissolved.
- 2. Phellonic acid $(C_{22}H_{43}O_8)$ has been isolated from cork, and this substance, together with its potassium salt, gives a red coloration with chlorzinc iodide. This suggests that the coloration of suberized membranes with chlorzinc iodide after

treatment with potash is due to the presence of potassium phellonate and not to cellulose, for, in addition, the coloration does not take place if the corky tissue be subjected to the action of boiling alcohol after treatment with potash.

3. After treatment with cuprammonia, the chlorzinc iodide gives a yellowish-brown colour; this, according to Gilson, is due to the alteration of potassium phellonate into the copper salt, and not to the removal of cellulose, as had been supposed.

Gilson separated from oak-cork suberic acid $(C_{17}H_{30}O_3)$ and phloionic acid $(C_{11}H_{21}O_4)$ in addition to phellonic acid. He does not think that these occur as true glycerine esters, since the suberin walls are insoluble in all fat-solvents, and do not melt at a temperature below 200° C.

These observations have been supported by Van Wisselingh,* who finds that the suberin constituents are mostly soluble in chloroform, and melt at a temperature below 100° C. He concludes that suberin consists of fatty substances with glyceryl or other compound esters easily decomposed by potash.

CONSTITUTION OF CELLULOSE.

The following characteristics of this substance throw some light on the constitution of cellulose:—

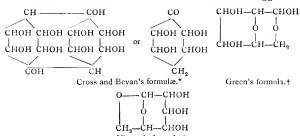
- 1. On hydrolysis it yields dextrose.
- On partial hydrolysis it sets free —CO groups which are present in some suppressed form in ordinary unchanged cellulose.
- 3. The fact that on destructive distillation it yields acetic acid and methyl alcohol points to the presence in the molecule of a $-CH_2-CO-$ grouping.
- 4. It is a very stable substance, which resists alkalis, oxidizing agents, and, to some extent, acetylation, except in the presence of a condensing agent, such as zinc chloride or sulphuric acid.
- 5. With strong acids it yields esters, e.g. nitrates, acetates, and benzoates.
 - 6. It undergoes the thio-carbonate reaction by treating

^{*} Van Wisselingh: "Arch. Neerland.," 1893, 26.

alkali cellulose with carbon disulphide, when a change, which may be represented by the following equation, takes place:—

$$RONa + CS_2 = CS \begin{cases} OR \\ SNa \end{cases}$$

Various alternative structural formulæ have been suggested.



Green § is of opinion that although cellulose is a colloid it does not follow that it has a high molecular weight. He considers that his formula well explains the following facts among others:—

- r. That the highest nitrate obtainable from a cellulose molecule containing six carbon atoms is a trinitrate.
- 2. That the highest acetate obtainable from a cellulose molecule containing six carbon atoms is a tri-acetate.
- 3. That cellulose does not react with phenylhydrazine, but on hydrolysis readily yields carbonyl groups which are able to react. The formula and its arguments are, however, not accepted by Cross and Bevan.¶

Dreaper regards cellulose as a typical colloid, and, as a

- * Cross and Bevan: "J. Chem. Soc.," 1901, 79, 366. + Green and Perkin: "J. Chem. Soc.," 1906, 81, 811.
- + Green and Perkin: "J. Chem. Soc.," 1906, 81, 811. ‡ Vignon: "Bull. Soc. Chim.," 1899, 21, 599.

§ Green: "Zeit. f. Farb. u. Textil Chemie," 1904, 3, 97, 309.

Il It should be clearly understood that by the nitration of cellulose it is possible to obtain a whole series of esters, representing different degrees of nitration. These various compounds may be described as mono-, di-, tri-, etc., up to deka, or possibly dodeka-nitrates of a cellulose molecule containing twenty-four carbon atoms. What is commonly called cellulose hexanitrate, the substance employed in the manufacture of gun-cotton is calculated on a C₁₂ molecule, which, therefore, corresponds to a trinitrate of a C₆ molecule.

¶ Cross and Bevan: "Zeit. f. Farb. u. Textil Chemie," 1904, 3, 197.

consequence, considers that it has no reacting unit such as a crystalline body has, nor has it a fixed molecular constitution such as can be represented by any constitutional formula; its reacting unit at any moment is a function of the condition under which it is placed.

INDUSTRIAL USES OF CELLULOSE AND CELLULOSE PRODUCTS.

One of the industries which consumes the largest amount of cellulose is that of paper manufacture. Formerly the chief sources of cellulose for this purpose were cotton or hemp fibres; but with the increased consumption of paper other sources had to be found. Although straw contains cellulose which has been only slightly lignified, it is found to be unsuitable for the preparation of pure cellulose, owing to the fact that it contains a considerable quantity of silica. The employment of wood as a source of cellulose became possible with the discovery of chemical methods of destroying the non-cellulose constituent lignin, i.e. the "encrusting substances," without affecting the cellulose proper.

In the manufacture of paper from linen rags or cotton waste the material is cut up, cleaned, and disintegrated by boiling successively with dilute sodium carbonate and caustic soda under pressure; the fibre is then bleached with chlorine, the excess being subsequently removed; it is then treated with resin, soap, and alum, and spread in thin layers and dried, whereby the fibres become felted together in a peculiar manner, with the formation of paper. When wood is used the "encrusting substances" may be removed by boiling with calcium bisulphite, whereby the lignin remains in solution and a fairly pure form of cellulose, known as sulphite cellulose, is produced. In the preparation of inferior quality papers the wood undergoes no chemical treatment for the removal of lignin; such papers can be recognized by applying to them any of the tests for lignocellulose. Cellulose used for the preparation of filter papers is, after the ordinary methods of purification, treated with hydrofluoric acid to remove silica.

COMMERCIALLY VALUABLE DERIVATIVES OF CELLULOSE.

When heated in a concentrated solution of zinc chloride, cellulose is converted into a viscid syrup. This syrup, when

forced through glass nozzles into alcohol, forms threads which, after being washed and carbonized, become hard and are used for electric lamp filaments; they have also been employed recently for the basis of incandescent lamp mantles.

Gun Cotton or Pyroxylin.—That a variety of different products may be obtained by the action of various strengths of nitric acid, either alone or in the presence of sulphuric acid, on cellulose, has already been mentioned. The substance known as gun cotton is a hexanitrate; it is obtained by immersing dry cotton waste, freed from grease by treatment with alkali. in a mixture of I part nitric acid (sp. gr. I:52) with 3 parts sulphuric acid (sp. gr. 1.84); the resulting substance is then rapidly and thoroughly washed with water, moulded into discs, and dried on heated plates. On explosion it produces corrosive gases and therefore is not suitable for use, as such. in firearms; when, however, the gun cotton is dissolved in ethyl acetate or acetone and the solution is evaporated, a new substance is obtained which has the same composition as gun cotton, but different properties; it explodes with less violence and produces no corrosive vapours, and is therefore employed in the manufacture of smokeless powder.

Blasting Gelatine is a mixture of gun cotton and nitroglycerine. Gun cotton mixed with a variety of other substances enters into the composition of numerous explosives, such as ballastite, melanite, cordite, etc. etc.

Collodion is the name applied to a solution of cellulose triand tetra-nitrates in a mixture of equal parts of 95 per cent alcohol and ether.

A substance known as artificial india-rubber* is produced by kneading together a mixture of tri- and tetra-nitrocelluloses partially dissolved in ether alcohol with castor oil. The resulting substance may be made to have any degree of elasticity, according to the materials which are mixed with it. It forms a more or less satisfactory substitute for rubber and possesses a high electric resistance. Though not explosive, it is inflammable, but to do away with this inconvenience the

^{*} This substance must be carefully distinguished from so-called Synthetic rubber, which is an artificially synthesized hydrocarbon of the formula $\{C_eH_g\}n$; this substance, if not actually identical with natural rubber, is at any rate closely related to it, whereas the artificial india-rubber mentioned above is a nitrated cellulose.

outer surface may be denitrated by treatment with alkali, whereby it is rendered non-flammable. *Artificial gutta-percha* is obtained by allowing an acetone solution of tetra-acetyl cellulose to evaporate.

Celluloid is produced by mixing the tri- and tetra-nitrates, as employed for collodion, with camphor.

Artificial Silks.—These are produced in a variety of ways by precipitating some form of cellulose from solution. The first artificial silk was prepared by Chardonnet, who obtained it by forcing collodion through fine nozzles; the thin stream of nitrocellulose solution on coming in contact with the air solidifies to a thread by the rapid evaporation of the solvent. To render it non-flammable the thread is denitrated by treatment with ammonium sulphide.

A second process for preparing artificial silk consists in dissolving bleached mercerized cotton (see p. 134) in cuprammonium solution. A fine stream of this solution is then run into a dilute sulphuric acid, whereby a continuous thread of cellulose is at once precipitated.

Viscose is obtained by acting on finely divided cellulose with soda and treating the resulting substances with carbon disulphide, whereby a cellulose thio-carbonate is produced; this substance on exposure to air decomposes spontaneously into cellulose alkali and carbon disulphide. Viscose solutions are employed for sizing paper, in the manufacture of wall-papers and for the production of thin threads for spinning by forcing the solution through fine nozzles and allowing the emerging stream to coagulate in the air.

Mixed with metallic dust and colouring matters, viscose can be converted into an artificial leather, and may also be employed for rendering canvas waterproof and for making cinematograph films, etc.

Viscoid, which is congealed viscose, is a hard mass obtained by mixing viscose with various substances and allowing the mixture to decompose spontaneously and harden; it is used for mouldings, cornices, statuettes, etc.*

Solid Spirit.—The substance sold under this name is obtained by pouring a solution of cellulose acetate in glacial

^{*}See Bersch: "Cellulose, Cellulose Products and Artificial Rubber," Philadelphia, 1904.

acetic acid into alcohol; a white solid is produced which does not melt, and burns when ignited without leaving any ash.

Finally, mention may be made of a few substances which are made from cellulose as a starting point, but which are produced only by the profound decomposition of the molecule. Thus by heating cellulose with a strong solution of caustic potash and soda, oxalic acid is produced, and by the destructive distillation of wood, acetic acid, acetone and methyl alcohol are obtained.

MICROCHEMICAL REACTIONS.

A. Normal Cellulose.

- With a dilute solution of iodine a yellow coloration results.
- After staining well with iodine, the addition of strong sulphuric acid causes the cellulose walls to swell considerably and to turn blue.
- 3. Chlorzinc iodide causes swelling, accompanied by the assumption of a blue colour.
- 4. Calcium chloride iodine solution turns pure cellulose rose-red, and finally violet.

Zimmermann gives the following directions for making this reagent. A concentrated solution of calcium chloride is made, and for each 10 c.c. of this solution there is added 5 gram of potassium iodide and 1 gram of iodine. The mixture is then gently heated and filtered through glass-wool.

- 5. Pure cellulose is easily soluble in cuprammonia.
- 6. The hemi-celluloses give different reactions; some turn blue with dilute iodine, and either do not dissolve in cuprammonia, or only after prolonged treatment.

B. Compound Celluloses.

(a) Lignin.

- 1. A brownish-yellow colour is given with iodine.
- 2. The addition of strong sulphuric acid, after previous treatment with iodine, turns lignified walls brown.
- 3. The same colour is obtained with the use of chlorzing iodide.

- 4. Calcium chloride iodine solution turns lignin yellow to yellow-brown.
 - 5. Insoluble in cuprammonia.
- 6. Aniline sulphate or aniline chloride in aqueous solution and acidified with the appropriate acid turns lignified walls a bright yellow.
- 7. If the sections be soaked for about a minute in an alcoholic solution of phloroglucin (or resorcin, hydroquinone, pyrogallol, or pyrrole) and then mounted in a drop of strong hydrochloric acid, the lignified walls are turned a bright red.
- 8. A concentrated solution of thallin sulphate in 50 per cent alcohol gives a yellow to orange-yellow coloration.

The sections should be treated first with alcohol, and the thallin sulphate solution should be freshly prepared.

The colour-reactions obtained by the use of aniline sulphate, thallin sulphate, phloroglucin and the other reagents mentioned in paragraph 7, are due to the presence of the furfurol complex in the lignin; any substance in the plant which contains this complex, e.g. coniferin, will give similar reactions.

- If lignified tissues be treated with chlorine water and then with sodium sulphide, a deep magenta colour is produced.
- 10. Lignocelluloses induce the formation of Prussian blue in the greenish-red solution produced by mixing ferric chloride with potassium ferricyanide.

(b) Suberin and Cutin.

- 1. With chlorzinc iodide, and also with iodine and sulphuric acid, a brown or yellow colour is given.
- 2. Suberized and cuticularized walls are insoluble in cuprammonia and concentrated sulphuric acid.
- 3. Suberized walls are coloured yellow with strong potash solution; on heating the colour deepens, and on boiling yellow oily drops exude from the membranes.
- 4. Suberized walls are the most resistant of membranes to Schultze's macerating mixture; but on boiling, oily drops of ceric acid are formed which are insoluble in carbon bisulphide but soluble in ether, benzol, and hot alcohol.
- 5. Suberized and cuticularized walls are stained green by the action of alcoholic solutions of chlorophyll. A strong

fresh solution of chlorophyll must be used, and the treatment should last for at least fifteen minutes in the dark. The sections may be washed in and examined in water. (Lignified walls are unacted upon by this and the following reagents.)

- 6. Similarly the same membranes are stained red by treatment with alcoholic solutions of Alkannin, Sudan III and Scharlach R.
- 7. If a section of the material be treated first with eau de Javelle, in order to destroy any tannins which may be present, suberized walls are stained very deeply with a solution of cyanin in 50 per cent alcohol to which an equal volume of glycerin has been added. Lignified walls will not be stained owing to the preliminary treatment with the eau de Javelle.
- 8. Corky walls are stained orange-yellow by an alcoholic solution of *extractum orleanæ* spirit which must be filtered before using.

FURTHER REFERENCES.

Cross and Bevan: "Researches on Cellulose," London, 1895, 1901, 1906, 1012.

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THE SYNTHESIS OF CARBOHYDRATES IN GREEN PLANTS.

INTRODUCTORY-ALDEHYDES.

In view of the important part played by aldehydes in questions relating to photosynthesis it appears desirable to draw attention to the chief properties of these substances before passing on to a consideration of the synthesis of carbohydrates in green plants.

It is, of course, well known that the aldehydes are the first products of the oxidation of primary alcohols:—

 $\begin{array}{c} CH_3OH + O = HCHO + H_2O\\ Methyl alcohol & Formaldehyde\\ CH_3CH_2OH + O = CH_3CHO + H_2O\\ Ethyl alcohol & Acetic aldehyde \end{array}$

The reconversion of formaldehyde into the alcohol can be effected by means of nascent hydrogen obtained by sodium amalgam and water.

Chemically, the aldehydes are very active, undergoing a number of reactions, some of which are of biological significance, whilst others serve as valuable means of isolation or identification.

- 1. Aldehydes are readily oxidized to the corresponding acids by even such mild oxidizing agents as ammoniacal silver hydroxide or Fehling's solution, or even atmospheric oxygen, as is shown by the following experiments:—
 - (a) A few drops of caustic potash are added to some silver nitrate solution in a test tube, ammonia is then carefully added, drop by drop, until the brown precipitate has just redissolved. A little dilute acetaldehyde solution is poured in and the mixture is warmed gently; if the solution be sufficiently dilute, a silver mirror will be deposited on the side of the test tube; otherwise a black precipitate will be formed:—

$$CH_3CHO + Ag_9O = CH_3COOH + 2Ag$$

(b) A little Fehling's solution is gently warmed with a few drops of dilute aldehyde solution; a change in colour takes place, from blue to green and yellow; finally the solution becomes colourless and a red precipitate of cuprous oxide (Cu_oO) comes down.

The readiness with which aldehydes are oxidized to acids accounts for the fact that most samples of aldehydes, unless freshly prepared, contain varying amounts of free acid.

2. Aldehydes are readily reduced by nascent hydrogen to the corresponding primary alcohols, according to the equation

$$CH_3CHO + 2H = CH_3CH_2OH$$

Acetic aldehyde Ethyl alcohol

- 3. Aldehydes restore the colour to Schiff's Reagent (a solution of magenta decolorised by sulphurous acid).
- 4. Aldehydes when warmed with caustic potash are converted into resinous substances of unknown composition. This can be readily shown with acetaldehyde; formaldehyde, however, when treated with potash undergoes a different change, being converted into a mixture of methyl alcohol and potassium formate, according to the equation

5. Aldehydes react with ammonia to form additive compounds; thus acetic aldehyde undergoes the following reaction:—

Here again formaldehyde behaves differently; if ammonia is added to a formaldehyde solution, it is neutralized quantitatively according to the equation

$$\begin{array}{ll} 6CH_2O + {}_4NH_3 = (CH_2)_6N_4 + 6H_2O \\ Formaldehyde & Hexamethylene \ tetramine \end{array}$$

with the formation of a crystalline solid which is used in medicine under the name of urotropine.

The reaction can be employed for estimating * the amount of formaldehyde in a solution by adding a known excess of standardized ammonia solution, and after some time titrating back the excess of ammonia by means of standard acid, using litmus as indicator.

Thus, for example, if 25 c.c. of the formaldehyde solution, after shaking with 50 c.c. of N/2 ammonia, required for neutralization 20 c.c. N/2 hydrochloric acid, then the amount of ammonia used up by the formaldehyde would be 50-20=30 c.c.

But 30 c.c. N/2 ammonia contain
$$\frac{30}{1000} \times \frac{17}{2} = .255$$
 gram NH_o

and since from the equation $4NH_{3}\left(68\right)$ are equivalent to $6CH_{2}O\left(180\right)$

.. 25 c.c. of the solution contained 0.68 gram formaldehyde.

6. With sodium bisulphite aldehydes form crystalline addition compounds which, being sparingly soluble in water, can be used for isolating aldehydes from mixtures.

Thus if some saturated sodium bisulphite solution be added to a fairly strong solution of aldehyde and the mixture shaken

For another method of estimating formaldehyde by weighing the mercury produced by the reduction of an alkaline solution of mercuric sulphite, see Feder: "Archiv d. Pharm.," 1907, 245, 25.

vigorously, a rise in temperature takes place accompanied by the formation of a white crystalline precipitate:—

 Aldehydes also form additive compounds with hydrogen cyanide; these compounds are known as hydroxycyanides or cyanohydrins:—

8. Aldehydes form crystalline compounds with hydroxylamine, phenylhydrazine, and semicarbazide; in all cases water is split off between the two reacting substances.

$$\begin{array}{l} CH_3CHO + NH_2OH = CH_3CH: NOH + H_2O \\ CH_3CHO + C_6H_5NHNH_2 = CH_3CH: N \cdot NHC_6H_5 + H_2O \end{array}$$

The resulting compounds, which are known as oximes, hydrazones or semi-carbazones, are usually substances with a characteristic crystalline form and melting point, which may be employed for the identification of the corresponding aldehydes. The use of phenylhydrazine for the identification of the sugars has already been described.

9. The aldehydes are able to react with alcohols with the formation of condensation compounds known as acetals; thus, for example, acetic aldehyde reacts with ethyl alcohol as follows:—

$$\begin{array}{c} \text{CH}_3 \\ \text{O} \\ \text{H} \\ \text{HOC}_2\text{H}_5 \\ \text{H} \\ \text{Acetic} \\ \text{aldehyde} \end{array} = \begin{array}{c} \text{CH}_3 \\ \text{OC}_2\text{H}_6 \\ \text{COC}_2\text{H}_5 \\ \text{H} \\ \text{Acetal} \\ \text{Acetal} \end{array}$$

By analogy, acetic aldehyde should also be able to react with water as follows:—

This substance does not, however, actually exist, since a compound having two or more hydroxyl groups attached to

the same carbon atom is, as a rule, unstable, and at once loses water. Exceptions to this rule are, however, occasionally met with; for example, chloral CCl₃CHO forms a stable compound, chloral hydrate, of the formula

[cf. Glucosides, p. 170].

10. Aldehydes exhibit a tendency to polymerize, that is, for two or more molecules to combine together to form new compounds of higher molecular weight.

Thus two molecules of formaldehyde will combine together, forming a compound known as paraformaldehyde (CH₂O)₂; this substance, which is a white solid, is obtained by evaporating an aqueous solution of formaldehyde.

A second polymer formed from three molecules of formal-dehyde is known as metaformaldehyde or trioxymethylene $(CH_2O)_3$. This substance is produced by the spontaneous polymerization of anhydrous formaldehyde.

In the case of both the above polymers the molecules of formaldehyde are probably connected together through oxygen atoms as under:—



which accounts for the fact that they are readily broken up into the simple molecules of formaldehyde by heating.

II. A different type of polymerization, involving the linking together of molecules of formaldehyde through carbon, is also known; this type of polymerization, which is sometimes known as aldol condensation, results in the formation of a more stable complex which cannot be reconverted into the simple substance.

The reaction takes its name from the substance produced by the action of dilute hydrochloric acid or zinc chloride on acetic aldehyde.

$CH_3CHO + CH_3CHO = CH_3CHOH.CH_2.CHO$ Aldol

The analogous reaction with formaldehyde is, however, brought about by dilute alkalis; in this way two molecules of formaldehyde give rise to glycollic aldehyde,

 $HCHO + HCHO = CH_2OH.CHO$ Glycollic aldehyde

or three molecules may combine together to produce glyceric aldehyde,

HCHO + HCHO + HCHO = CH₂OH.CHOH.CHO Glyceric aldehyde

By repeatedly shaking a 4 per cent solution of formal-dehyde for half an hour with an excess of lime water, and then filtering the solution and setting it aside for some days until the odour of formaldehyde had disappeared, Loew * was able to obtain a crude mixture of sugars called formose, from which true reducing hexose sugars have been isolated. This change may be represented by the equation:—

$$6HCHO = C_6H_{12}O_6$$

Similarly H. and A. Euler † have shown that when a 2 per cent solution of formaldehyde is heated for some hours with calcium carbonate, a pentose sugar—arabinoketose—is produced; in addition to this substance, glycollic aldehyde and dihydroxyacetone are produced, but in smaller quantity.

FORMALDEHYDE.

From the point of view of photosynthesis formaldehyde is of outstanding interest; as is well known, it is at ordinary temperatures a colourless gas with a pungent odour; when cooled to -21° it condenses to a liquid. It is usually met with in the form of an aqueous solution, commercial formalin, which contains about 40 per cent of the gas dissolved in water and is used as a disinfectant or as a hardening medium for pathological and other specimens and occasionally as a preservative for milk. It undergoes most of the general reactions for aldehydes which have been mentioned above.

^{*}Loew: "Ber. deut. chem. Gesells.," 1887, 20, 142, 3039; 1881, 21, 270; 1889, 22, 470, 478.

⁺ Euler, H, and A.: id., 1906, 39, 36, 39.

Its peculiar behaviour towards ammonia, resulting in the formation of hexamethylene tetramine, has already been mentioned; this substance, which is used under the name of urotropine, is a crystalline base which dissolves in hot or cold water; with bromine it forms an additive compound—tetrabromo-hexamethylene tetramine $(CH_2)_8N_4Br_4$ —which has been used for detecting small quantities of formaldehyde in solution (see p. 153).

Formaldehyde also reacts with ammonium salts as well as with free ammonia, as follows:—

This reaction has been made use of as a means of estimating ammonium salts in solution by titrating the amount of free acid liberated according to the above equation on adding sufficient formaldehyde to a solution containing ammonium salts. For this purpose both the formaldehyde solution and the solution to be analysed must be previously neutralized, if necessary. An excess of the neutralized formaldehyde solution is then added to a known volume of the solution containing the ammonium salts, and after thoroughly shaking for one or two minutes the amount of acid set free is determined by titration with standard caustic soda, using methyl orange as indicator; the amount of ammonia can be calculated from the fact that each 36.5 grams of hydrochloric acid liberated correspond to 17 grams of ammonia.

The reactions most suitable for characterizing small quantities of formaldehyde are as follows:—

Rimini's test consists in adding 2 drops of phenylhydrazine hydrochloride, 2 drops of sodium nitroprusside solution, and I c.c. of sodium hydroxide to I c.c. of the liquid to be tested. A blue colour is formed, which changes rapidly through green and brown to red. Schryver* has modified this test and made it much more sensitive; he recommends the following method: to IO c.c. of the liquid to be tested add 2 c.c. of a I per cent solution of phenylhydrazine hydrochloride freshly made up and filtered; then add I c.c. of a 5 per cent solution of sodium ferricyanide, also freshly made up, and 5 c.c. of hydrochloric acid; a brilliant magenta colour is

^{*} Schryver: " Proc. Roy. Soc., Lond.," B., 1910, 82, 226.

produced. The test is a very delicate one and will detect quantities of formaldehyde varying from I part in I,000,000 to I part in I00,000. Acetic aldehyde gives no colour with this reagent.

The following test, due to Denigés,* is sensitive for formal-dehyde, even in presence of acetic aldehyde up to 2 per cent; 5 c.c. of an aqueous solution of formaldehyde are mixed with 1'2 c.c. of pure sulphuric acid (sp. gr. 1'66) and 5 c.c. of Schiff's reagent. An intense violet colour having an absorption band in the orange is produced. Schiff's reagent may be prepared by adding a litre of 0'01 per cent of solution of magenta to 20 c.c. of sodium hydrogen sulphite solution (sp. gr. 1'3), and after five minutes adding 20 c.c. of hydrochloric acid (sp. gr. 1'18).

Kimpflin† tested for formaldehyde in the leaf of Agave mexicana by injecting into it, by means of a capillary tube, a concentrated solution of sodium hydrogen sulphite, containing an excess of pmethylamino-mcresol. The presence of formaldehyde was indicated by the formation of a red precipitate on exposure to light. The precipitate is best seen by examining a section of the leaf which has been dipped in absolute alcohol. Formaldehyde is the only aldehyde giving a stable red colour with the above reagent, but other aldehydes give unstable green, yellow, or reddish-brown colours.

In 1906, Usher and Priestly ‡ stated that they obtained formaldehyde by subjecting green leaves to steam distillation; the presence of formaldehyde in the distillate was proved by evaporating it down with ammonia and adding bromine water to the residue to convert the hexamethylene tetramine so formed into its tetrabromo derivative.

In the opinion of Curtius and Franzen, § the presence of formaldehyde in plants has not been established with certainty by previous workers, inasmuch as the majority of the above tests for the substance in question are inconclusive, since they are also given by other aldehydes. By working with 1500 kilos of hornbeam leaves these authors conclusively proved the

^{*} Denigés: "Compt. rend.," 1910, 150, 529. † Kimpflin: "Compt. rend.," 1907, 144, 148.

Usher and Priestly: "Proc. Roy. Soc., Lond.," B., 1906, 77, 369.

[§] Curtius and Franzen: "Ber. deut. chem. Gesells.," 1912, 45, 1715.

presence of formaldehyde by oxidizing it to formic acid, which was recognized by the usual tests.

A critical review of the investigations regarding the presence of formaldehyde in plants has been published by Polacci.*

PHOTOSYNTHESIS.

The formation of carbohydrates takes place in living chlorophyll-containing cells on exposure to sunlight: the initial substances are the carbon dioxide of the atmosphere and the water contained within the cells; the end products are oxygen and carbohydrate, the volume of oxygen evolved being practically equal to the volume of carbon dioxide used up; the processes take place with extraordinary rapidity, oxygen being evolved almost immediately after exposure to light, and the carbohydrate appearing very quickly afterwards.

This rapidity of action means that, whatever the intermediate products may be, they must be very transient, so that small amounts rather than large quantities are to be looked for in healthy leaves which are actively engaged in photosynthesis. It appears from the investigations of Friedel† that the quantity of oxygen of the air is immaterial to the process. The amount of this oxygen may be decreased to 2 per cent or increased to 50 per cent without interfering with the photosynthetic processes.

In the laboratory it is convenient to take the presence of starch as an indicator of the fact of photosynthesis in microscopic experiments; but the first recognizable products of photolysis are soluble carbohydrates, which may, later on, be converted into starch, as in the potato, bean, etc., or may remain as sugar, as in Allium, Scilla, and many other plants. In these latter, starch may be formed provided the sugar in the leaves be allowed to become concentrated.

In addition to sugars, other compounds such as mannite, oils, tannin, and organic acids have also been described as direct products of photosynthesis; whether they have been so described on sufficient grounds or not, their occurrence in

^{*} Polacci: "Atti. R. Acc. Lincei," 1907, [v.], 16, 1. 199. + Friedel: "U.S. Dept. Agric.," 1901, Bull, 28.

this connexion is rare enough to justify their exclusion from the present consideration.

The above facts are well established, but with regard to the intermediate products, and to the nature of the processes concerned, our knowledge is as yet very incomplete.

In 1870 Baeyer put forward the hypothesis that the carbon dioxide is split up by the plant into carbon monoxide and oxygen, and that the water is concurrently resolved into its constituent elements. The carbon monoxide and hydrogen thus produced then combine to produce formaldehyde, which undergoes polymerization, and so forms a hexose.

These changes may be represented in the following equations:—

$$\begin{cases} \text{1. } CO_2 = CO + O \\ \text{2. } H_2O = H_2 + O \\ \text{3. } CO + H_2 = CH_2O \\ \text{4. } 6(CH_2O) = C_3H_{12}O_6 \end{cases}$$

Thus, according to the theory, there are two distinct actions; the first leading to the formation of formaldehyde, and the second to the production of sugar.

Considering the first part of Baeyer's theory, it is seen that both carbon monoxide and hydrogen are supposed to be produced, but carbon monoxide has not been found in a free state in the living plant, nor is it a substance which lends itself at all readily to constructive metabolism, the evidence as to whether plants are able to make use of it for synthetic purposes being contradictory. Bottomley and Jackson* state that if the carbon dioxide normally present in the atmosphere be replaced by about twenty times as much carbon monoxide—the increase in the amount of the latter being necessary on account of its lesser solubility in water as compared with carbon dioxide—plants of Tropwolum formed starch and flourished. Preliminary analyses also showed that, in the case of seeds germinated in an atmosphere in which the carbon dioxide had been replaced by carbon monoxide, there was in the seedlings an increase in organic carbon. Further, they found that a negative pressure obtained in the vessels containing the plants assimilating carbon monoxide. This was to be expected if the hypothesis be accepted, for if the

^{*} Bottomley and Jackson: "Proc. Roy. Soc., Lond.," B., 1903, 72, 130.

carbon monoxide be used up in photosynthesis, then the amount of oxygen set free would be half that evolved in normal photolysis. On the other hand Kraschénnikoff* has come to the conclusion, based on a number of experiments, that green plants cannot make use of carbon monoxide; he points out, however, that his evidence does not prove that carbon monoxide is not formed in the early stages of photosynthesis. It may also be remarked that according to the investigations of Sulander,† carbon monoxide acts as an anæsthetic, but is much weaker in its action than chloroform. He found that '5 per cent of this gas was sufficient to inhibit the growth of seedlings of the lupin and the germination of the spores of many Fungi. Of course it does not therefore follow that carbon monoxide is not formed in plants. As is well known, carbon dioxide is itself an anæsthetic if present in a sufficient amount, and possibly it is more potent in this respect than is carbon monoxide, for Seelander found that in many cases the streaming movements of protoplasm were not affected even after several hours' exposure to the last-named gas.

A modification of Baeyer's theory thus appears to be necessary. Erlenmeyer, long before the experimental work on carbon monoxide just referred to was done, suggested that the carbonic acid in the cells undergoes a reduction which leads to the formation of formic acid and oxygen, and that the formic acid is further reduced to formaldehyde and oxygen:—

1.
$$H_2CO_3 = CH_2O_2 + O$$

2. $CH_2O_2 = CH_2O + O$

or else that the action is continuous and that the carbon dioxide and water may directly give rise to formaldehyde and oxygen:—

$$CO_2 + H_2O = CH_2O + O_2$$

According to these views either formaldehyde or formic acid must be produced.

These substances, as is well known, are poisonous, so that, if formed, they must be polymerized before they have time to injure the protoplasm, and experiments have shown that

^{*} Kraschénnikoff: "Rev. Gen. Bot.," 1909, 21, 177. † Sulander: "Beih. bot. Centrbl.," 1909, 24, I, 357.

under certain conditions, formaldehyde may be made use of by the plant. Thus Bokorny* showed that *Spirogyra* can make starch when supplied with a compound of formaldehyde and sodium hydrogen sulphite; also Treboux† and Bouilhac have stated that *Elodea*, *Sinapis*, and certain Algæ can form starch in the dark when supplied with dilute ('0005 per cent) solutions of formaldehyde.

In this connexion Grafe's ‡ results are important, for he found that if green seedlings be grown in the light in an atmosphere containing no carbon dioxide but formaldehyde vapour (not more than 1.3 per cent), they show a greater increase of growth and in dry weight as compared with controls grown under the same conditions, but without the formaldehyde.

The next question which naturally arises is whether formaldehyde occurs in assimilating leaves, and whether it is possible to reproduce *in vitro*, with the aid of a suitable sensitizer, the preliminary action which is supposed to take place in the plant, viz., the formation of formaldehyde from water and carbon dioxide

Reinke § in 1883 was one of the first instigators to discover formaldehyde in green leaves, and Curtius and Reinke some years later stated that aldehydes occur in chlorophyll-containing cells, provided they be exposed to light; these substances, however, do not occur in Fungi. Amongst the first to attempt to reproduce in a test tube the supposed initial photosynthetic stages was Bach, who states that formaldehyde is produced from carbon dioxide in the presence of water by the action of sunlight, provided that a suitable optical sensitizer, such as uranium acetate or dimethyl aniline, be employed. In other words, the formation of formaldehyde in the leaf is not a vital process. He also found that this same substance is produced from carbonic acid in the presence of hydrogen palladium, which acts as the reducing agent.

^{*}Bokorny: "Biol. Centrbl.," 1897, 17, 1; "Ber. deut. chem. Gesells.," 1891, 24, 103. †Treboux: "Flora," 1903, 92, 49.

[†] Ireboux. Piota, 1903, 92, 49. ‡ Grafe: "Ber. deut. bot. Gesells.," 1911, 29, 19. § Reinke: "Ber. deut. bot. Gesells.," 1883, 1, 406.

^{||} Curtius and Reinke: "Ber. deut. chem. Gesells.," 1897, 30, 201.

Polacci* as a result of his observations came to the conclusion that formaldehyde is of constant occurrence in green leaves; also that the carbonic acid is reduced by hydrogen, which is given off by the plant, to formaldehyde, water and oxvgen.

According to Friedel, the presence of living organisms is not essential in this synthetic process, as indicated by the evolution of oxygen and absorption of carbon dioxide. process takes place by means of an enzyme which he obtained from the leaves of the spinach by extraction with glycerine. This extract is mixed with alcohol sufficient in amount to precipitate the enzyme, which is then filtered off and dissolved in water. Leaves of the same plant are dried at 100° C., and powdered up. If a mixture of the powder and the extract be exposed to light, carbon dioxide will be absorbed and oxygen evolved.

Macchiati + obtained similar results, and further found that the leaf powder alone would give off oxygen when suspended in water and exposed to sunlight.

On the other hand Bernard t and others from similar experiments obtained no positive results.

Beyerinck, by using different methods, also found that leaf powder would give off oxygen in the presence of light.

In this particular connexion, the work of Molisch § is of much importance. He found that the extracted sap from fresh leaves can give off oxygen in the presence of sunlight. and the same result is obtained by using the leaves of Lamium album dried at 35° C. and exhibiting no signs of life. If, however, these extracts be filtered through a fine porcelain candle, no such positive results are obtained, and further, the unfiltered extracts lose their power after boiling.

In considering these results Blackman | points out that in all experiments designed to show extra-cellular photosynthetic processes, in which powdered leaves or leaf extracts are used, it must be borne in mind that the results may be due to the

^{*} Polacci: "Atti. Inst. Bot. Pavia," 1900, 6; 1902, 8; 1904, 10.

⁺ Macchiati: " Rev. Gen. Bot.," 1903, 15, 20. # Bernard: "Beih. bot. Centrbl.," 1904, 16, 36.

[§] Molisch: "Bot. Zeit.," 1904, 62, 1.

[|] Blackman: "New Phytol.," 1904, 3, 33.

residual vitality of expressed protoplasm, and not to a relatively simple enzyme, as is sometimes supposed; and in reviewing the position, he states that "the work of Molisch carries with it the greatest conviction, and leads one to conclude that photosynthesis can exist to a small degree apart from the living cell. One may further hazard the hypothesis that this function is correlated with some machinery more complex than an enzyme, but much less complex than a complete protoplasmic unit."

The work of Bach * and Polacci has been adversely criticized by Euler.+ Polacci found that if an extract of active leaves be distilled, the presence of formaldehyde may be detected in the distillate. On repeating these experiments, Euler found that the extract gave a feebler reaction when compared with the distillate and, what is of greater importance, ordinary hay gave results as good as those obtained from fresh green leaves. Euler also experimentally examined Bach's results; it will be remembered that Bach found that carbon dioxide and water may be made to combine in the presence of sunlight, provided that a sensitizer, not necessarily chlorophyll, be present. Euler found that if carbon dioxide be not used, formaldehyde is still produced, although in smaller quantities; further, that the passage of nitrogen or hydrogen, in place of the carbon dioxide, through the solution of uranium acetate, yielded as good results; finally, in the experiments of which dimethylaniline was used, the resulting formaldehyde was due to impurities in the sensitizer employed.

The question has received renewed attention recently owing to the conclusions arrived at by Usher and Priestly.‡ Summarizing their results, they found that normally the photolysis of carbon dioxide and water leads to the formation of hydrogen peroxide and formaldehyde, although under certain conditions formic acid may be produced. Of these two products, the hydrogen peroxide is decomposed by an enzyme in the plant. and the formaldehyde is condensed by the protoplasm. Thus in the presence of light and chlorophyll carbon dioxide and

^{*} Bach: "Compt. rend.," 1893, 116, 1145. + Euler: "Ber. deut. chem. Gesells.," 1904, 37, 3411.

Lusher and Priestly: "Proc. Roy. Soc., Lond.," B., 1906, 77, 369; 1906, 78, 318; 1911, 84, 101.

water give rise to hydrogen peroxide and formaldehyde; the former is acted upon by an enzyme so that oxygen is given off; the latter by the action of the protoplasm is polymerized into carbohydrate. If the formaldehyde be not used up rapidly it poisons the enzyme, so that the peroxide of hydrogen is not decomposed, and will thus destroy the chlorophyll. They also found that the photolysis of carbon diox de may take place in vitro, provided one of the products be removed.

From further experiments they conclude that the formation of formaldehyde from carbonic acid in the presence of chlorophyll is independent of vital or of enzymic activity; the products of such decomposition are formaldehyde and peroxide of hydrogen, formic acid being an intermediate product. Also they find that it is possible to reconstruct the photosynthetic process outside the green plant as far as the production of formaldehyde and oxygen, by the aid of a suitable catalysing enzyme, and as far as the production of oxygen and starch by the aid of non-chlorophyllous protoplasm together with the enzyme.

They, in addition, repeated Bach's experiments referred to above and found that formic acid is a product of the photolysis of carbonic acid in the presence of an inorganic salt of uranium; and that formaldehyde is probably formed as a transitory intermediate product.

In the following year, 1907, Fenton * found that in the presence of magnesium—which substance, there is reason to suppose, is the active principle of chlorophyll—formaldehyde may be obtained from an aqueous solution of carbon dioxide, more especially if weak bases be present.

Usher and Priestly also found that an aqueous solution of carbon dioxide could be decomposed by the a and β rays from radium emanation. The action of '0001 c.c. of radium emanation on 200 c.c. of water saturated with carbon dioxide resulted in four weeks in the production of hydrogen peroxide and formaldehyde. Most of the latter was in a polymerized state, but the solution contained no sugar.

Similar results were obtained by the action, on solutions of carbon dioxide, of the ultra violet rays given off by a quartz

^{*} Fenton: "J. Chem. Soc., Lond.," 1907, 91, 687.

mercury-vapour lamp, so that it appears that "ultra-violet light can effect a measurable decomposition of aqueous carbon dioxide without the intervention of an optical or chemical sensitizer, whilst under normal conditions some such agent is required; moreover, the results furnish very strong support for the belief that both formaldehyde and hydrogen peroxide are formed in a green leaf".

The following experiments, also by Usher and Priestly, point to the same conclusion:—

A glass plate was covered with a layer of gelatine made up with a solution of catalase. When set, the film was painted over with a film of chlorophyll, and the plate sealed up in a glass tube which contained air and some caustic potash. The tube was set up at 5 p.m., and at noon the next day the chlorophyll was quite green and distorted with bubbles. In a control experiment, lacking the catalase, the chlorophyll showed signs of bleaching after one hour, and at the end of the experiment was much bleached.

Two other experiments were then set up (11.30 p.m.); one exactly as the first, and the other having a solution of carbon dioxide in place of the potash. At 11 a.m. the next day the chlorophyll of the last was considerably bleached and showed the presence of relatively much formaldehyde, whilst the chlorophyll of the former was quite green, and only slight indications of aldehyde were obtained.

As a result of such experiments Usher and Priestly have no doubt that "the bleaching of chlorophyll in sunlight, whether carbon dioxide is present or not, is due to the formation of hydrogen peroxide. . . . As regards the production of formaldehyde, the experiments are equally conclusive in showing that it is only detected by Schiff's reagent when carbon dioxide is present."

Using the living plant, the results were of the same nature, but were not so consistent, owing to the difficulty of controlling the experiments.

With regard to the evolution of oxygen, they confirm the work of Molisch on the evolution of oxygen on the killed green leaves of *Lamium*. Usher and Priestly used *Elodea* which had been anæsthetized for fifteen hours, and found that in the presence of carbon dioxide, oxygen is evolved in the

light. They also describe an experiment to show that a parallel result is obtained in vitro.

A Petri dish is divided into two by a strip of cork cemented across the middle. A culture of luminous bacteria which are extraordinarily sensitive to free oxygen and glow only in its presence—in nutrient gelatine was poured into one compartment, and in the other was placed the same culture but with catalase added. When the gelatine was set, a film of chlorophyll was painted over the surface. The lid was then put on and firmly fixed with wax and gold size in order to make the joint quite air-tight. The preparation was then placed in a dark room. After the lapse of two days no glow was discernible. The dish was then exposed to daylight for five minutes, and once more examined by the observer in the dark room. The bacteria were found to be glowing, the side containing no catalase showing the most light, a difference which at first sight is surprising, but it was found that the bacteria would glow in the presence of very small quantities of hydrogen peroxide.

There is another point of considerable importance. The formation of formaldehyde and hydrogen peroxide from carbonic acid involves the absorption of heat, so that a film of chlorophyll placed in an atmosphere containing moist carbon dioxide should, when illuminated, be at a lower temperature than a control film in an atmosphere free from carbon dioxide. By the use of a thermo-junction and a suspended coil galvanometer this was found to be the case.

Some of the earlier experiments and conclusions arrived at by Usher and Priestly have been adversely criticized by Blackman * and Ewart +; their later work, however, places their conclusions on a much firmer basis.

Ewart's conclusion that chlorophyll contains formaldehyde in a combined state has been confirmed by Schryver,‡ who has published certain observations on the subject. He finds that formaldehyde is more abundant in chlorophyll films after exposure to bright sunlight than when exposed to a dull light. He also states that if glass plates covered with films of

^{*} Blackman: "New Phytol.," 1906, 5, 132. † Ewart: "Proc. Roy. Soc., Lond.," B., 1908, 80, 30. ‡ Schryver: "Proc. Roy. Soc., Lond.," B., 1910, 82, 226. See also the further work of Wager, and Warner (p. 230).

chlorophyll be kept in the dark no formaldehyde is produced, no matter whether moist carbon dioxide be present or not: further, if such plates be exposed to sunlight in an atmosphere free from carbon dioxide, a very minute quantity of formaldehyde is produced: on the other hand, the plates when exposed to the sun's rays in the presence of moist carbon dioxide give a distinct formaldehyde reaction.

From this Schryver concludes that in the presence of sunlight, water, and carbon dioxide, there is a continuous production of formaldehyde, which is continually being condensed to sugar. If this condensation does not proceed rapidly enough to remove all the formaldehyde, the excess enters into combination with the chlorophyll, and, as the free formaldehyde is used up, this compound of formaldehyde with chlorophyll decomposes, setting free the former, which is converted into sugar. There is thus in the plant a mechanism by means of which the quantity of free formaldehyde is regulated, so that at no time is the amount sufficiently large to become toxic; thus Curtius and Franzen found only '0008613 gram of formaldehyde per kilo of green hornbeam leaves (see p. 153).

With regard to questions relating to the form of energy used in these photosynthetic processes, a few passing remarks may be made. As is well known, the view commonly held is that radiant energy, more especially those rays of the red end of the spectrum, afforded by the sun, is the direct source. And as this is adequately treated in most texts on plant-physiology, it is not proposed to deal with it here. At the same time it is to be borne in mind that other forms of energy may be made use of by the plant.

Royer * brought about the electrolytic reduction of carbon dioxide, and by similar means Coehn,+ in 1904, produced formic acid from this same compound. Brodie found that by means of a silent discharge formaldehyde, together with marsh gas, was produced from a mixture of hydrogen and carbon dioxide; and Löb, in 1906, found that formaldehyde may be produced by the action of a silent discharge of elec-

^{*} Royer: "Compt. rend.," 1870, 70, 731.

[†] Coehn: "Ber. deut. chem. Gesells.," 1904, 34, 2836, 3593. ‡ Brodie: "Proc. Roy. Soc., Lond.," 1874, 22, 171.

[§] Löb: "Zeit. Electrochem.," 1906, 12, 282.

tricity through a solution of carbon dioxide in water. Fenton * also has pointed out that the synthetic action of light and of the silent electrical discharge are practically identical. Thus there is evidence which suggests that electric energy may play a part in the earlier processes of photosynthesis; a suggestion which is supported by the fact that, according to Polacci,† the formation of carbohydrates is promoted in leaves by electrical energy, provided it be not too intense, especially when a continuous current is made to pass directly into the tissues.

As a result of a number of experiments, Gibson ‡ comes to the conclusion that the light rays which are absorbed by the chlorophyll are transformed into electrical energy, and it is this transformed energy which brings about the decomposition of carbonic acid to formaldehyde and oxygen.

With regard to other forms of energy, the results obtained by Usher and Priestly with radium emanations and with ultra violet rays have been mentioned. Attention may be drawn also to the work of Kernbaum, \S who found that water exposed to the influence of β rays and of ultra-violet rays led to the production of hydrogen and hydrogen peroxide. Also Berthelot and Gaudechon $\|$ found that formaldehyde is produced by the action of ultra-violet rays on carbon dioxide in the presence of a reducing agent.

The question now arises as to how the sugars are produced from the formaldehyde, presuming the latter to be formed. Living leaves contain many different sugars; thus Meyer, in 1885, found that cane sugar, maltose, dextrose and levulose all were present. Later, Brown and Morris \P described that, in the case of the leaves of Tropxolum, cane sugar accumulated while the hexoses remained practically constant in amount. Meyer also states that the di- and polysaccharides increase while the photosynthetic processes are active, while, in the dark, the monosaccharides increase.

If Baeyer's hypothesis be correct, then it would naturally

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* Fenton: "J. Chem. Soc., Lond.," 1907, 91, 687.

† Polacci: "Atti. Inst. Bot., Pavia," 1905, II, II, 7.

‡ Gibson: "Ann. Bot.," 1908, 22. 117.

§ Kernbaum: "Compt. rend.," 1909, 148, 705; 1909, 149, 273.

# Berthelot and Gaudechon: id., 1910, 150, 1690.

¶ Brown and Morris: "J. Chem. Soc., Lond.," 1893, 63, 604.
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be expected that during light, provided always the other requisite conditions obtained, the hexoses would increase in amount. So far, this has not been satisfactorily proven. It is to be borne in mind, however, that a portion of these hexoses, in all probability, would be used for the immediate requirements of the plant: thus a certain proportion would be immediately used up in respiration; another part might possibly be accounted for as being necessary for the synthesis of proteins; again, a certain quantity would be translocated to other parts of the plant; finally, it must not be forgotten that the whole process of photosynthesis takes place with extraordinary rapidity, starch, a final product, often appearing directly after the chloroplast is exposed to light.

Considering first the formation of a simple sugar, it is possible, of course, that the formaldehyde may at once undergo polymerization into a polyose sugar; but this can only be assumed when all intermediate products have been proved to be absent. And, as regards these intermediate products, although our present physiological knowledge is decidedly scanty, indeed, practically non-existent, there are certain laboratory facts which are of the greatest importance.

As already mentioned above, the molecular formula of a hexose shows that it is a polymer of formaldehyde. The first successful attempt to bring about such a polymerization was made by Butlerow in 1861, who, by the catalytic action of lime water, at the ordinary temperatures, on trioxymethylene (itself a polymer of formaldehyde), obtained a syrup with a somewhat bitter taste, which he called methylenitan. Subsequently Loew undertook an investigation of the action of milk of lime on formaldehyde, and devised the following experiment. A four per cent solution of formaldehyde is mixed with an excess of milk of lime and repeatedly shaken for about half an hour; after filtering, the mixture is set aside for some days until the pungent smell of formaldehyde has disappeared. The solution, which will now reduce Fehling's solution, yields a colourless syrup with a very sweet taste. This substance, which is known as crude formose, was examined by Emil Fischer, who found it to consist of a mixture of various hexoses and succeeded in isolating from it a small quantity of a sugar—acrose. This same sugar can also be obtained by the

action of dilute caustic soda on a substance called glycerose, which is obtained by the oxidation of glycerol. Similarly arabino-ketose, a pentose which has not yet been identified in plants, has been synthesized by the action of calcium carbonate, a very mild catalysing agent, from formaldehyde. From the acrose thus obtained, Fischer was able by an elaborate series of reactions to prepare ordinary fructose or levulose.

It is thus seen that the conversion of formaldehyde to fructose is reasonably complete, and when it has been proven beyond all shadow of doubt that formaldehyde can be formed from carbon dioxide and water by a photosynthetic method, the chain of chemical evidence will be complete. In order to bring the vital process into line with the laboratory process, it is necessary to prove the presence of the intermediate products in the chlorenchyma of plants, together with suitable catalysts.

With regard to the formation of the higher carbohydrates from the fructose, practically nothing is known, and unfortunately our present knowledge of the constitution of disaccharides is limited.

While it is a very easy matter to determine what two sugars are obtained by the hydrolysis of any given disaccharide, it is as yet merely a matter of speculation as to how these two sugars are united in the disaccharide molecule. In the case of cane sugar, for example, it is known that it yields on hydrolysis a mixture of dextrose and levulose, and in attempting to assign a structural formula to cane sugar it must be borne in mind that the substance has no longer the properties of either an aldehyde or a ketone,* and therefore the formula must be without the characteristic —CHO or —CO groups.

The formation of dextrose does not present so great a difficulty, for fructose has been converted into dextrose, in vitro; it is only necessary to find a suitable agent in the green tissues of plants to bring about the isomerization. Thus, having the dextrose and fructose, the sucrose could be produced by their condensation. But at the same time it must be remarked that the synthesis of disaccharides from monosaccharides has been achieved only in a very few cases,

^{*} Cane sugar does not react with phenylhydrazine, and does not reduce Fehling's solution.

and all attempts to synthesize cane sugar from glucose and fructose, or from invert sugar, have hitherto failed. Fischer and Armstrong* were able to synthesize a disaccharide isolactose-by the action of an enzyme, Kefir lactase, on a mixture of glucose and galactose; the same authors also synthesized melibiose. Similarly isomaltose has been obtained by Croft-Hill † by the action of maltase on glucose.

The cane sugar is often supposed to be assimilated by the protoplasm, which in turn forms the starch; finally the maltose may be formed from the starch, or by the condensation of dextrose which has already been accomplished in the laboratory.

In conclusion, mention should be made of a photosynthesis of carbohydrate in the absence of chlorophyll which was effected by Stoklasa and Zdobnicky.‡ Light from a quartz mercury lamp was allowed to pass through a mica window into a vessel containing a mixture of carbon dioxide and hydrogen; formaldehyde was slowly produced and this, in presence of caustic potash, was polymerized with formation of a sugar or mixture of sugars which was optically inactive and not fermentable by yeast. The authors suggest that the chlorophyll in plants acts as a means of absorbing ultra-violet ravs.

According to Löb, however, these conclusions are not justified by the experiments performed by Stoklasa and Zdobnicky.

With regard to the reverse process, Bertholet and Gaudechon || found that carbohydrates are decomposed by sunlight and by ultra-violet light from a mercury lamp. The products of decomposition are carbon monoxide, carbon dioxide, methane and hydrogen; aldehydic sugars differ from ketonic sugars both in the readiness with which they are decomposed and in the composition of the gaseous mixtures produced.

^{*} Fischer and Armstrong: "Ber. deut. chem. Gesells.," 1902, 35, 3144.

⁺ Croft-Hill: "J. Chem. Soc., Lond.," 1898, 73, 634; see also Emmerling: "Ber. deut. chem. Gesells.," 1901, 34, 600, 2206.

‡ Stoklasa and Zdobnicky: "Chem. Zeit.," 1910, 945.

[&]amp; Löb: "Biochem. Zeitschr.," 1912, 43, 434.

^{||} Bertholet and Gaudechon: "Compt. rend.," 1910, 151, 395; 1912, 155, 401, 831.

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SECTION III.

GLUCOSIDES.

THE glucosides are compounds of some complexity which on decomposition yield glucose together with one or more other substances, usually of an aromatic nature; they are, therefore, often described as ether-like compounds of carbohydrates with aromatic compounds. The carbohydrate in the majority of cases is glucose, but occasionally it may be an isomeric hexose, such as galactose or even a pentose such as rhamnose. Thus, for example, digitonin, one of the glucosides of Digitulis purpurea, yields on hydrolysis both glucose and galactose, while hesperidin and quercitrin, the glucosides contained respectively in the unripe orange and in the bark of Quercus tinctoria, give rhamnose.

The reaction by means of which glucosides are split up into their constituent parts is in almost all cases one of hydrolysis,* and can, therefore, usually be brought about by boiling with dilute mineral acids or, in some cases, alkalis. In nature, however, the decomposition is effected by means of suitable ferments which often exist in the same part, although generally in different cells to the glucoside.

Each glucoside may have an enzyme appropriate to itself, but any one particular ferment may have the power of splitting several glucosides.

Thus, for example, the glucoside amygdalin is hydrolysed by its appropriate enzyme emulsin to glucose, benzaldehyde and hydrocyanic acid, according to the equation:—

$$C_{20}H_{27}NO_{11} + 2H_2O = 2C_6H_{12}O_6 + C_6H_5CHO + HCN$$

^{*}The decomposition of potassium myronate into potassium hydrogen sulphate, glucose and allyl mustard oil can hardly be described as hydrolysis (see p. 186).

whilst the enzyme amygdalase,* contained in yeast, is only able to effect a partial hydrolysis to glucose and mandelonitrile glucoside.†

$$C_{20}H_{27}NO_{11} + H_2O = C_6H_{12}O_6 + C_{14}H_{17}NO_6$$

On the other hand, emulsin can hydrolyse other glucosides, besides amygdalin, such as salicin, arbutin, etc.

THE CONSTITUTION OF THE GLUCOSIDES.

The constitution of the natural glucosides can be best understood by a brief consideration of the simplest known artificial glucosides which have been synthesized from glucose.

The constitution of glucose is ordinarily represented by the formula CH₂OH CHOH CHOH CHOH CHOH CHOH, which shows it to be a pentahydric alcohol and an aldehyde at the same time. When dissolved in water, however, it behaves in a peculiar manner, exhibiting the phenomenon of muta-rotation, that is to say, the optical activity of the resulting solution does not attain a steady value until some time after the solution has been made up.

The change is supposed to be connected with some alteration in its molecular configuration which may be explained by assuming that the compound

is temporarily formed, † but that water is thereupon split off again between one of the hydroxyl groups of the terminal carbon atom and the hydroxyl attached to the fourth carbon atom as follows:—

^{*} Caldwell and Courtauld: "Proc. Roy. Soc., Lond.," B., 1907, 79, 350.

⁺Fischer: "Ber. deut. chem. Gesells.," 1899, 28, 1509.

[‡] Compare the formation of similar compounds from other aldehydes (p. 149).

This latter formula for glucose, showing the asymmetry of the terminal carbon atom, marked with an asterisk, explains the possibility of the existence of two optically isomeric forms of glucose,* and also accounts for the ability of glucose to react with methyl alcohol to form two isomeric α - and β -methyl glucosides † according to the equation:—

A number of analogous compounds have since been prepared by Fischer and his co-workers from mannose, galactose, and fructose, the resulting compounds being termed mannosides, galactosides, and fructosides respectively.[‡]

As the result of studying the action of the two enzymes, maltase and emulsin, upon other glucosides, Fischer divides these substances into two classes known as a-glucosides and β -glucosides, of which maltase can only split the a-glucosides and emulsin the β -glucosides. Amygdalin is, therefore, an a- β -glucoside, e.g. of its two glucose molecules, one is the a- and the other is the β -modification.

Since most natural glucosides yield on hydrolysis a substance containing a hydroxyl group, it seems reasonable to assume that the original glucoside was formed by a reaction similar to the one given for the artificial glucosides mentioned above, that is by the elimination of water between a hydroxyl group of the sugar and one from the other compound. On this assumption the constitution of some of the better known natural glucosides could be represented as follows:—

^{*}Tanret: "Compt. rend.," 1895, 120, 1060; cf. also Fischer: "Ber. deut. chem. Gesells.," 1893, 26, 2400; Lowry: "J. Chem. Soc, Lond.," 1899, 75, 213; and Perkin: "J. Chem. Soc, Lond.," 1902, 81, 177.

⁺ The specific nature of enzymic hydrolysis is exhibited in the case of these two artificially synthesized α - and β -methyl glucosides which Fischer prepared by the action of hydrochloric acid on a solution of glucose in methyl alcohol. The α -glucoside, which is dextro-rotatory, is hydrolysed by maltase, but not by emulsin, while the β -glucoside, on the contrary, is unaffected by maltase, but is hydrolysed by emulsin (see also section on Enzymes, p. 353).

[‡] Fischer and Fischer: "Ber. deut. chem. Gesells.," 1910, 43, 2521.

Moreover, the constitution of amygdalin itself would appear to be best represented by the formula *:—

and, by analogy, phaseolunatin is regarded as the glucose ether of acetone cyanhydrin (see p. 182).

In some cases the natural glucosides have been actually synthesized; thus salicin has been obtained by the reduction of the corresponding aldehyde glucoside, helicin:—

$$C_6H_{11}O_5 \cdot O \cdot C_6H_4CHO + 2H = C_6H_{11}O_5 \cdot O \cdot C_6H_4CH_2OH$$

the helicin itself having been synthesized from glucose and

salicylic aldehyde.

Identification.

For the identification of glucosides the character of the cleavage products are relied upon; these products, with the

exception of sugar, are frequently of a volatile nature and possess a characteristic smell; their chemical nature, however, varies so much that there is no general test for the whole group other than that afforded by the presence of a carbohydrate after hydrolysis.

The method devised by Bourquelot has been much employed; the expressed sap of the plant is examined as to its power of reducing Fehling's solution and as to its optical properties, the ferment emulsin is then added to the extract and the mixture kept in the warm for a time. The amount of sugar is thereupon again estimated and so also is its rotatory power. The increase indicates roughly the amount of glucoside originally present.

In microchemical work the same test, in a simplified form, may be applied, the preparation being treated with a solution of emulsin or of dilute sulphuric acid, and then gently warmed with Fehling's solution.

These tests, however, are of value only in certain cases, for some glucosides, and for that matter some other substances likewise, can reduce Fehling's solution without undergoing a preliminary hydrolysis, also there may be present in the cells of the plant other substances which on hydrolysis would yield glucose, and further some glucosides like gynocardin exhibit a remarkable resistance to the action of acids.

The decomposition products of many glucosides are brightly coloured, for example, rhinanthin when boiled with dilute hydrochloric acid yields a dark blue-green colour which is due to rhinanthogenin. Several give a brilliant red coloration with strong sulphuric acid, e.g. salicin and phloridzin. Most glucosides are soluble in varying degrees in either hot or cold water and alcohol; the majority are insoluble, or nearly so, in ether, which fact is made use of in separating them from alcoholic solutions.

PHYSIOLOGICAL SIGNIFICANCE OF GLUCOSIDES.

In attempting to assign the part played by these substances in the economy of the plant, it must be remembered that the number of glucosides of natural occurrence are very numerous and, in some cases, of a diverse nature; it is, therefore, possible that the significance of the presence of one glucoside may be quite different to that of another, but even in the case of glucosides of the same nature there is much diversity of opinion. They have been described, on insufficient grounds, as direct products of photosynthesis. Many consider them to be of value as food-stuffs on account of the sugar they contain; the occurrence of certain glucosides in seeds lends some support to this view, for in the case of the bitter almond hydrocyanic acid, in the free state, may be identified when germination starts, also the observations of Treub,* who found that in the case of some plants containing cyanogenetic glucosides the amount of the latter decreased if the plant was placed in the dark, in order that photosynthesis could not take place. On the other hand there was an increase in quantity when the plants were exposed to light, and this increase reached a maximum at about midday.

Weevers† considers that salicin, populin, arbutin and similar glucosides are of the nature of reserve food-materials, for not only is the formation of these substances a suitable means for the storage of sugar on account of their low diffusibility, but the facts of their seasonal or diurnal variation lend support to this opinion. Thus in Vaccinium Vitis-Idea the arbutin is stored in the leaves, and when the new leaves are formed in the spring it is used up; it is split by a suitable enzyme, the sugar is used up, and the hydroquinone remains behind and combines with more sugar, so that by the autumn the leaves once more contain much arbutin.

In the case of the willow, salicin is formed day by day, but during the night it is split by salicase into sugar and the alcohol saligenin. The glucose is translocated, and the saligenin remains behind and is converted into salicin by combining with sugar the next day. This process stops in the autumn, by which time there is relatively much salicin in the cortex of the stem.

This translocation of glucosides from the leaves of many plants—but not of all, *Sambucus* and *Indigofera* being exceptions—is significant, and so also are the facts relating to the

^{*} Treub: "Ann. Jard. Bot. Buitenzorg," 1896, 13, 1; 1907, 21, 79, 107; 1910, 23, 85.

⁺Weevers: "Kon. Akad. Wet.," Amsterdam, 1902; "Rec. Trav. Bot. Néerl," 1910, 7, 1.

amount of glucosides in the bark and other parts of plants at different seasons of the year. Thus in Salix and Populus the glucoside (salicin) is most abundant in the autumn and winter, and is used up in the following spring during the period of flowering and seed formation; also in the case of Taxus the glucoside (taxicatin), which appears principally in the young shoots, is greatest in amount in the autumn and winter. In Pangium edule and other plants the amount of cyanogenetic glucosides is greatest in young leaves, with increasing age the amount diminishes.

Guignard* does not believe that glucosides, or at any rate the cyanogenetic ones, are reserve food-stuffs, and the fact that, if introduced into the food materials of a plant, glucosides have an injurious effect, owing to the aromatic residues, gives some support to this contention.

A contrary opinion is held by Peche,† who holds that hydrocyanic acid is a direct product of photosynthesis; some of it combines with sugar to form a glucoside, and some is transported in a labile form, probably in a loose combination with tannin, and stored for future use as food in various tissues.

The occurrence of certain glucosides, especially in places of active metabolism such as leaves and young shoots, may indicate that certain bye-products are fixed, either temporarily or more permanently, in this form.

In conclusion it may be stated that many may perform a biological function; thus the bitterness or poisonous nature of the glucosides or of the products of hydrolysis, other than sugar, may serve as a protection against herbivorous or fruiteating animals; the antiseptic properties of these dissociation products may have a value in preventing the development of disease organisms in parts which may be damaged, e.g. seeds, leaves and bark. Some may play a part in connexion with the secretion of sugar by extrafloral nectaries, for it appears that the basal cells of these structures, together with the elements of the adjacent tissues, are rich in glucosides.

^{*}Guignard: "Compt. rend.," 1905, 141, 236; 1906, 143, 451. †Peche: "Sitz. Kais. Akad., Vienna," 1912, 121, 33.

CYANOGENETIC GLUCOSIDES.

Among the more important glucosides are the cyanogenetic ones, so named because on hydrolysis they yield hydrocyanic acid as one of the products.

Hydrocyanic acid is of fairly common occurrence in the higher plants, and although sometimes it occurs in the free state it is, in the majority of cases, combined; the nature of many of these compounds has not yet been ascertained, but it is not improbable that generally they are glucosides.

Cyanogenetic glucosides, although widely distributed, are somewhat rare when compared with other glucosides such as the saponins. Hydrocyanic acid has been found in a few Fungi, and in certain plants of the following Natural Orders of the higher plants: Polypodiaceæ, Aroideæ, Gramineæ, Sapindaceæ, Sapotaceæ, Proteaceæ, Ranunculaceæ, Magnoliaceæ, Lauraceæ, Droseraceæ, Rosaceæ, Saxifragaceæ, Leguminosæ, Platanaceæ, Euphorbiaceæ, Compositæ, etc. It will be observed from this list that some Cohorts, for example Rosales and Ranales, stand out in having several natural orders characterized by the presence of the substance in question.

In the individual plant the cyanogenetic glucosides occur more especially in the leaves and buds, in the seed, and also in the bark.

In Pangium edule Treub* found such glucosides in the phloem, pericycle, and in special cells of the leaves; Guignard† describes such compounds as occurring in the leaves of vigorous shoots, the young bark, and in the unripe fruit of Sambucus nigra and species of Ribes. The amount present in a member is not constant; Verschaffelt‡ found that as the buds of Prunus Padus and P. Laurocerasus open, the amount of hydrocyanic compounds increases as rapidly as do the other substances present. Treub has found that in plants growing in the tropics and which contain cyanogenetic glucosides, these substances disappear before leaf-fall; in some cases this depletion is quite sudden, in others the glucosides gradually

^{*} Treub: "Ann. Jard. Bot. Buitenzorg," 1907, 21, 107.

[†] Loc. cit.

[‡]Verschaffelt: "Kon. Akad. Weten. Amsterdam," 1902.

disappear. On the other hand in *Indigofera* and *Sambucus* the glucosides are not removed before the fall of the leaves.

Treub also states that the amount present depends on the quantity of available sugar; he observed that there obtains a daily variation, the maximum quantity occurring at about mid-day. It has also been ascertained that the quantity of cyanogenetic glucosides in Pangium, Phaseolus lunatus, Zea and Sorghum may be increased by the application of manures, rich in nitrates; on the other hand, it must be pointed out that in some cases, e.g. Phaseolus lunatus, the glucoside may be eliminated from the seed by suitable methods of cultivation. In some examples of seeds which contain little or no hydrocyanic acid there may be a marked increase on germination, thus in the flax, Dunstan and Henry* found that the seeds contained '008 per cent of the acid, whereas in the seedlings '135 per cent obtained; the same increase also occurs in the sweet almond

These authors also point out that the percentage of hydrocyanic acid in *Linum*, *Sorghum*, *Lotus arabicus* and *Zea Mais* gradually increases to a maximum and then decreases, sometimes to zero.

The stage of development at which the maximum is reached varies in the different plants; thus, to take two extreme cases, in the flax the maximum obtains when the seedlings are between four and five inches high, whilst in *Lotus arabicus* the maximum occurs at the period of flowering.

From these observations it is clear that the actual amount of the substance in question varies pretty considerably; it may be very small or relatively large, thus in the young leaves of *Pangium* the presence of 3 per cent of hydrocyanic acid has been ascertained. In this connexion attention may be drawn to observations of Greshoff,† who states that the absence of hydrocyanic acid does not necessarily indicate the absence of a cyanogenetic glucoside. If the glucoside produced be very small in amount, as in the case of *Xeranthemum*, the hydro-

Dunstan and Henry: "Brit. Assoc. Rep., York," 1906; "Phil. Trans. Roy. Soc., Lond.," 1907, B., 194, 515; "Proc. Roy. Soc., Lond.," B., 1900, 67, 224; 1901, 68, 374; 1903, 72, 285.

[†]Greshoff: "British Assoc. Rep., York," 1906, 138; "Kew Bull.," 1909, 397.

cyanic acid may be used up directly it is formed, so that benzaldehyde only will be found as a decomposition product.

ISOLATION OF CYANOGENETIC GLUCOSIDES.

Dunstan and Henry give the following method for the isolation of dhurrin from Sorghum vulgare. The plants are dried at a low temperature and ground up as finely as possible. The material so obtained is extracted with alcohol and filtered; the alcohol is then distilled off from the filtrate and the residue dissolved as completely as possible in warm water. Lead acetate is added to this aqueous solution until no more precipitate (chiefly lead tannate) comes down. A current of sulphuretted hydrogen—a large excess is to be avoided—is then passed through the filtrate and the lead sulphide filtered off. The excess of sulphuretted hydrogen can be removed from the filtrate by passing through it a current of air. liquid is then worked up with pure animal charcoal, sufficient in amount to convert the whole, when dry, into a powder, and dried in a vacuum desiccator. When quite dry the material is extracted with anhydrous ethyl acetate in a Soxhlet apparatus; this solvent slowly removes the glucoside, leaving most of the sugar and other impurities behind. On distilling off the solvent a syrup remains which may, if necessary, be again treated in the same fashion. The syrup will deposit crystals of the glucoside after standing for a few days in a vacuum over sulphuric acid. The crystals so obtained may be recrystallized from hot alcohol or boiling water.

CHEMISTRY OF CYANOGENETIC GLUCOSIDES.

We may now pass on to a brief consideration of the chemical nature of the cyanogenetic glucosides. These glucosides vary in different plants. Thus comparing the cherry laurel, Prumus Laurocerasus, with Pangium edule, it has been found that in the former plant the localization of the glucosides is not so clearly defined as in the latter; also this substance disappears from the leaves of the cherry laurel, when kept in the dark, much more slowly than does the glucoside in Pangium on similar treatment. This indicates that these two glucosides have a different chemical constitution, and analysis has shown this to be the case. In Pangium edule, and also in Linum

and other plants, the glucoside has an acetone cyanhydrin residue, while in the case of *Prunus* the residue is benzaldehyde cyanhydrin. The former glucosides are less stable than the latter.

With regard to the stages which lead up to the formation of prussic acid and its compounds, Gautier has put forward the supposition that it may possibly be formed by the action of formaldehyde on nitrates, and this view is not inconsistent with the distribution of nitrates in the leaves of some plants, but nothing definite is known.

Reactions, Microchemical and Otherwise.

- I. The presence of cyanogenetic glucosides or of free hydrocyanic acid can generally be detected by chewing a small piece of the material.
- 2. Thoroughly crush the part it is desired to examine under water and set it aside for some time, then filter and add to the filtrate a little silver nitrate; a white precipitate indicates hydrocyanic acid, but this test must be used with caution as many other substances give a white precipitate with silver nitrate.

If the amount of enzyme present in the tissue be very small, the maceration must be allowed to proceed for some time, or emulsin may be added to hasten the decomposition.

3. Cut a thick section of the fresh tissue to be examined and place it in a 5 per cent alcoholic solution of potash for about a minute; transfer to a solution containing 2.5 per cent ferrous sulphate and 1 per cent ferric chloride and keep at about 60° C. for ten minutes. Place the preparation in a dilute solution of hydrochloric acid—one part of strong acid to six parts of water—for five to fifteen minutes. The presence of hydrocyanic acid is indicated by the formation of Prussian blue.

If leaves are to be tested, instead of cutting them up they may be pricked all over with a bunch of fine needles and then treated as above.

4. Guignard's Test.—Dip strips of white filter-paper in a 1 per cent solution of picric acid and dry, moisten the papers again with a 10 per cent solution of sodium carbonate and

again dry. The test papers, which may be kept in stoppered bottles for some time without deterioration, turn an orange red in the presence of fumes of hydrocyanic acid. The test is very delicate, and the rapidity of the change in colour depends on the amount of prussic acid present, so that if the quantity be very small the paper may have to be suspended in the test tube containing the material to be tested, for a day.

This test has been modified by Waller so as to give quantitative results, but it has been pointed out by Chapman* that the coloration is due to reduction, and is, therefore, not specific for hydrocyanic acid; accordingly the method must be used with caution.

It was found that if a leaf of the cherry laurel, *Prunus Laurocerasus*, be immersed in an aqueous solution containing of per cent picric acid and 5 per cent sodium carbonate, the leaf is unharmed and the fluid undergoes no obvious change. If, however, the leaf be immersed in the same fluid to which chloroform has been added in the proportion of 4 c.c. per 100 c.c. of fluid, the formation of hydrocyanic acid takes place and the sodium picrate turns red. The intensity of the colour is the basis of the quantitative estimation of the hydrocyanic acid.

The standard colour is obtained by mixing together equal volumes of the picrate fluid and '002 per cent hydrocyanic acid. This mixture, which contains 10 mgs. of hydrocyanic acid per litre, is allowed to stand for twenty-four hours in an incubator kept at 40° C. The intensity of the colour is designated by the symbol T10, and corresponds to 10 mgs. of hydrocyanic acid per litre; a colour intensity of T1 similarly corresponds to 1 mg. of hydrocyanic acid per litre. By diluting the standard solution, a gamut of colour intensities may be obtained T1, T2, T3 . . . the figure in each case corresponding to the number of milligrams of hydrocyanic acid in one litre of fluid. These colours may be matched closely by aqueous solutions of potassium bichromate.

In making up the standard solution it is important to allow the mixture to stand in an incubator kept at a temperature of 40° C. for certainly not less than an hour owing to the slowness of development of the full tint.

^{*} Chapman: "Analyst," 1910, 35, 469. See also Francis and Connell: "J. Amer. Chem. Soc.," 1913, 35, 1629.

AMYGDALIN.

Amygdalin, $C_{20}H_{27}NO_{11}$, is a lævo-rotatory bitter substance which is fairly soluble in water, and gives with concentrated sulphuric acid a pale reddish-violet coloration, this, however, is not a distinctive test, since the same coloration is given by other glucosides, e.g. menyanthin.

Amygdalin occurs in the seeds of the bitter almond, Pyrus Amygdalus; it is, however, generally stated not to occur in the seeds of the cultivated almond, the sweet variety, although emulsin, its appropriate enzyme, is present. Dunstan and Henry have shown that traces of hydrocyanic acid occur in the seeds, and more than traces in the seedlings, of the sweet almond; it is probable, therefore, that a small quantity of amygdalin does occur in the sweet variety. This relative absence of glucoside in the cultivated plant is important, and the same phenomenon has been found to obtain, by these same authors, in Phaseolus lunatus. The seeds of the wild plant yield large quantities of hydrocyanic acid, whereas those of the cultivated plants give very little or none.

Amygdalin has also been described as occurring in *Pyrus Malus*, *Pyrus Aucuparia*, *Pyrus cydonia* and other plants.

This glucoside is generally obtained by crushing the seeds of the bitter almond and subjecting the mass to considerable pressure between hot iron plates in order to remove the oil. The solid cake is then digested with hot alcohol which dissolves out the amygdalin. The alcoholic extract is evaporated down when the amygdalin separates out in scale-like crystals belonging to the monoclinic system.

It has already been mentioned that the appropriate enzyme generally occurs in the same tissues as the glucoside; this being so, the bitter almonds have only to be crushed in water in order to bring the ferment emulsin into contact with the amygdalin to bring about the hydrolysis.

This change probably takes place in two stages:-

I.
$$C_{20}H_{27}NO_{11} + H_2O = C_6H_{12}O_6 + C_{14}H_{12}NO_6$$

Mandelonitrile glucoside
II. $C_{14}H_{12}NO_6 + H_2O = C_6H_{12}O_6 + HCN + C_6H_6CHO$

If crystals of amygdalin be dissolved in water and then subjected to the action of maltase the hydrolysis will not proceed further than is represented in the first of the two above equations; emulsin, on the other hand, can hydrolyse the mandelonitrile glucoside as indicated in the second equation, and, of course, it can bring about the whole series of changes.

The formation of mandelonitrile glucoside in this process is of some interest since it is isomeric with the glucoside sambunigrin which occurs in the fruit of the elder, *Sambucus nigra*.

The crude oil of bitter almonds contains hydrocyanic acid which may be removed by distillation with lime and ferrous chloride which converts the prussic acid into Prussian blue. Pure benzaldehyde is a colourless or pale yellow liquid, soluble in alcohol, but practically insoluble in water. Its specific gravity is 1.05, and its boiling point 180° C. On exposure to air it becomes converted into benzoic acid.

DHURRIN.

This is a glucose closely allied to amygdalin, and occurs in the seedlings of *Sorghum vulgare*, but not in the older plants; it has the empirical formula $C_{14}H_{17}NO_7$ and yields, on hydrolysis, glucose, hydrocyanic acid and parahydroxybenzaldehyde:—

$$C_{14}H_{17}NO_7 + H_2O = C_6H_{12}O_6 + HCN + C_6H_4OHCHO$$

Similar glucosides occur in the seedlings of Panicum and $Z\epsilon a$.

PHASEOLUNATIN.

Phaseolunatin, $C_{10}H_{17}O_6N$, occurs in the seeds of wild plants of *Phaseolus lunatus*, it is present only in very small quantities, or is entirely absent from the seeds of the cultivated plants. It is also present in *Linum* and many rubberyielding plants, such as *Hevea braziliensis* and species of *Manihot*. Associated with it in its natural surroundings is the enzyme phaseolunatase which is able to hydrolyse it to acetone, glucose, and prussic acid.*

^{*} Dunstan, Henry and Auld: "Proc. Roy. Soc., Lond.," B., 1906, 78, 145, 152.

LOTUSIN.

Lotusin, $C_{28}H_{31}NO_{16}$, occurs in *Lotus arabicus*. It is a bitter, yellow-coloured substance, which when fresh does not reduce Fehling's solution.

On hydrolysis it yields glucose, hydrocyanic acid, and lotoflavin, a bright yellow crystalline precipitate:—

$$C_{29}H_{31}NO_{16} + 2H_2O = 2C_6H_{12}O_6 + HCN + C_{15}H_{10}O_6$$

Lotoflavin

Lotusin, like dhurrin, does not occur in old plants with ripe seeds, it is present only in the younger stages of development.

It is hardly necessary to point out the economic importance of this occurrence of cyanogenetic glucosides in the younger stages of plants like *Lotus arabicus* and *Sorghum*; much loss of stock has been sustained by their consumption by cattle.

SAPONINS.

According to the researches of Greshoff,* the saponins are very widely distributed in the higher plants; he has identified them in various plants belonging to the natural orders: Piperaceæ, Saururaceæ, Primulaceæ, Loganiaceæ, Oleaceæ, Polemoniaceæ, Proteaceæ, Caprifoliaceæ, Compositæ, Cucurbitaceæ, the majority of the natural orders of the cohort Centrospermæ, Ranunculaceæ, Magnoliaceæ, Leguminosæ, Rosaceæ, Saxifragaceæ, Pittosporaceæ, Polygalaceæ, Rutaceæ, Rhamnaceæ, Guttiferæ, Thymelæaceæ, Combretaceæ, Myrtaceæ, Lecythidaceæ, Araliaceæ, Gramineæ, Liliaceæ, and Gleicheniaceæ.

The term saponin, though originally restricted to a specific substance obtained from the root of *Saponaria rubra* and *S. alba*, is now applied to a large group of compounds, all of which have properties similar to those possessed by the original saponin.

General Properties and Uses of Saponins.

The saponins are mostly amorphous colloidal substances which dissolve readily in water; their aqueous solutions, if shaken up alone, produce a froth, but if shaken in the presence

^{*} Greshoff; "Kew Bulletin," 1909, 397.

of oils, fats or resins, they produce emulsions which are characterized by their great stability.

Connected with their emulsifying property is the employment of saponins as substitutes for soaps, a fact which is indicated in the name Saponin itself and also by the names Saponaria, soap wort and Quillaia (meaning wash wood), etc.

The so-called soap nuts are the fruits of Sapindus (fructus saponis indici) and these, as well as the beans of Entada scandens and Lychnis chalcedonica or Tartary soap, are largely used in the East for washing clothes, since they have no deleterious effect on the colour or the fibre of the most delicate fabrics.

Aqueous solutions of saponins have a marked power of retaining dissolved gases, as, for example, carbon dioxide; for this reason saponins are occasionally added to effervescent drinks, such as ginger-beer or lemonade, a use which is to be deprecated owing to their toxic properties.*

Occasionally saponins are employed for making suspensions of solids in water since they exert an inhibiting effect on the precipitation or deposition of suspended solids. Concentrated aqueous solutions of the saponins have adhesive properties.

Solubility.

The saponins are, as a rule, neutral substances which dissolve readily in water, but a few are acid in character and require a small quantity of alkali to enable them to dissolve completely.

For their aqueous solutions saponins may be precipitated unchanged by the addition of ammonium sulphate.

In the form of lead or barium compounds they may be precipitated from aqueous solutions by the addition of either lead acetate or basic acetate of lead or by means of a solution of barium hydrate.

The saponins are almost all insoluble in absolute alcohol, ether, chloroform and benzene.

^{*}The saponin obtained from the bark and wood of Guajacum officinale is occasionally used for this purpose since it is practically non-poisonous, its hæmolytic action (see p. 185) being only very slight.

Physiological Action.

The saponins are characterized by their strongly marked toxic properties. Fishes, particularly, are very sensitive to saponins, being killed by a solution of one part in 100,000 parts of water, a fact which is made use of by fishermen in the East, as the fish killed by these means are not unfit for human consumption.

Saponins have a powerful solvent action on blood corpuscles, a property which is known as hæmolysis. This property may be connected with their tendency to combine with cholesterol,* which substance they abstract from the blood corpuscles thereby rendering them soluble.

The action may be illustrated by adding a small quantity of a solution of a saponin † in 0.9 per cent sodium chloride to I c.c. of a solution made by dissolving I c.c. of defibrinated blood in 100 c.c. of 0.9 per cent sodium chloride; after a short time the blood corpuscles will have dissolved leaving a clear solution.

The hæmolytic action may be destroyed by shaking up some of the saponin solution with an ethereal solution of cholesterol and then warming for some hours at 36° C.; this treatment causes the saponin to combine with cholesterol to produce an inactive compound which has no solvent action on blood corpuscles.

Chemistry of the Saponins.

As already stated, the majority of saponins are neutral substances, while a few have feebly acid properties. Only a single saponin, namely, Solanin, has basic properties; this substance, which occurs in *Solanum nigrum*, *S. dulcanura* and in the fruit of potatoes, owes its basic property to the presence of a nitrogen atom (see Nitrogen Bases, p. 263), and appears to form a connecting link between the saponins and the alkaloids

The neutral saponins are precipitated from solution by basic lead acetate, while acid saponins are precipitated by lead

^{*} They also combine with phytosterol.

[†]The saponins of Guajacum officinale and Bulnesia Sarmienti have hardly any hamolytic action, and hence are only slightly toxic,

acetate. Similarly, barium hydroxide precipitates neutral saponins in the form of their barium compounds (see below).

On hydrolysis with dilute mineral acids* the saponins yield sugars such as glucose, galactose, arabinose, and rhamnose, together with other substances termed sapogenins, the constitution of which is unknown.

The nature of the sapogenin obtained from any particular saponin varies with the conditions of the hydrolysis; in some cases careful hydrolysis may yield a primary sapogenin and a sugar, while more complete hydrolysis gives rise to an end sapogenin together with more sugar.

The hydrolysis of Digitonin, the saponin contained in *Digitalis purpurea*, may, according to Kiliani, be represented by the equation:—

$$\begin{array}{c} C_{54}H_{92}O_{28} + \, _2H_2O = C_{30}H_{48}O_6 + \, _2C_6H_{12}O_6 + \, _2C_6H_{12}O_6 \\ Digitonin & Digitogenin \ Glucose & Galactose \end{array}$$

On mixing together alcoholic solutions of a saponin and of cholesterol a precipitate of the cholesterol compound is at once formed. These cholesterol compounds are, as a rule, easily decomposed; in most cases, prolonged extraction with ether will remove the cholesterol, and the saponin is recovered unchanged and possesses its original physiological action.

The saponins are reducing agents, and will reduce ammoniacal silver nitrate to metallic silver; similarly, prolonged boiling with mercuric chloride reduces this substance to calomel; saponins also blue a solution of potassium ferricyanide containing ferric chloride, by reducing the ferric salt to the ferrous condition, and so giving rise to the formation of Turnbull's blue.

If boiled with acetic anhydride, alone or in presence of sodium acetate or zinc chloride, the saponins are converted into acetyl derivatives which are no longer toxic. On boiling the acetyl derivatives with alcoholic potash the acetyl groups are removed, but the resulting compound is not identical with the original saponin.

When treated with a hot saturated solution of baryta a

^{*} Hydrolysis can, in some cases, be effected by bacteria, and Quillaia saponin is even said to be hydrolysed by emulsin (see Gonnermann: "Pflüger's Archiv," 1906, 113, 185).

saponin is precipitated in the form of a barium compound. If this latter is treated with the requisite amount of sulphuric acid the barium may be completely removed, but the resulting substance, unlike the original saponin, is physiologically inactive.

Reactions.

The following reactions are made use of in demonstrating the presence of a saponin:—

- I. Aqueous extracts readily form a froth when shaken up.
- 2. Concentrated sulphuric acid gives with all saponins, either in the cold or on warming, a violet or red colour.
- 3. Concentrated sulphuric acid containing a little ferric chloride gives with many saponins a blue or bluish-green colour or fluorescence.
- 4. The hæmolytic action described on page 185 may be tried.

Although the above reactions are best carried out in the test tube, numbers 2 and 3 may be made use of in microchemical work.

OTHER GLUCOSIDES.

In addition to the above, there occur in plants a large number of other glucosides which do not readily lend themselves to reasonable classifications. The exigencies of space will permit of reference only to the following, which are among the more important and more interesting of them.

SINIGRIN.

Sinigrin, or myronate of potash, occurs in the seeds of certain Cruciferæ, notably Sinapis nigra.

Preparation.

Green gives the following method for its extraction: One kilogram of the seeds of the black mustard is ground to a fine powder, and then extracted with one and a half litres of 82 per cent alcohol. The mixture is heated on a water bath until the volume of the alcohol is reduced by about one-sixth, the alcoholic extract is then filtered off, and the residue pressed

while it is still hot. The operation is then repeated; the residue is dried at 100° C., and digested for twelve hours with eight times its volume of cold water. A small quantity of barium carbonate is added to this aqueous extract, which is then evaporated to a syrup. The sinigrin is contained in this syrup, and is extracted by boiling with 82 per cent alcohol. Finally, the alcoholic extract is evaporated down, when the glucoside crystallizes out in rhombic prisms, which are freely soluble in water and warm alcohol, but much less soluble in cold

Sinigrin is split by the enzyme myrosin into glucose, potassium hydrogen sulphate and allyl isothiocyanate, or mustard oil, which may be recognized by its distinctive smell.

$$C_{10}H_{18}O_{10}NKS_2 = C_6H_{12}O_6 + KHSO_4 + CH_2: CHCH_2NCS$$

CONIFERIN.

This glucoside occurs in various coniferous trees, especially in young parenchyma, and also in asparagus. With concentrated sulphuric acid coniferin gives a violet coloration, while hydrochloric acid and phenol give a blue coloration. In brief, most reagents used in the demonstration of the lignification of cell walls (p. 145) give similar reactions both with coniferin and vanillin, and for this reason it is supposed that both these substances occur in such thickened walls. The use of these reagents tends to show that coniferin is more abundant in young wood cells, whilst vanillin, which is coloured yellow by thallin, occurs more extensively in the older elements. All such colour reactions, however, must be used with caution since many of them depend on the presence of certain complexes, e.g. aldehyde, which may occur in the molecules of widely different substances.

Coniferin crystallizes in needle-shaped crystals, m.p. 185°, and is soluble in warm water and warm alcohol. On hydrolysis by mineral acids or by emulsin it gives glucose and coniferyl alcohol:—

$$C_{16}H_{22}O_8 + H_2O = C_6H_{12}O_6 + C_{10}H_{12}O_3$$

Coniferin Coniferyi

The latter is a crystalline substance melting at 73°.

Both coniferin and coniferyl alcohol when oxidized with

potassium bichromate and sulphuric acid yield vanillin, the aromatic constituent of the fruits of Vanilla planifolia.

The reaction was formerly employed for the preparation of artificial vanillin, but has now been replaced by the oxidation of isoeugenol, which is obtained by the action of dilute alkalis upon eugenol, a substance contained in oil of cloves.

The relationship between these three substances is as follows:—

SALICIN.

Salicin, $C_{13}H_{18}O_7$, occurs in the bark of *Salix viminalis*. It has a bitter taste and crystallizes in colourless prisms and scales. It is sparingly soluble in cold water but is more soluble in hot alcohol, especially amyl alcohol, and may be extracted from aqueous solutions by means of this solvent. Microscopically, salicin is indicated by the fact that it gives a bright red colour with strong sulphuric acid, also with Fröhde's reagent * it yields a violet coloration. On steeping the section in a solution of emulsin, saligenin is produced which gives a blue colour with ferric chloride.

Preparation.

Salicin may be prepared by boiling the willow bark with water which will extract a certain amount of tannin, colouring and other matters together with the salicin. The greater part of impurities may be precipitated by the addition of lead acetate. The precipitate is then filtered and a stream of sulphuretted hydrogen is passed through the filtrate in order to remove the lead. The filtrate on evaporation yields crystals of salicin which may be further purified by recrystallization from alcohol.

Another method is to treat the bark with benzene in order

^{*} Sodium molybdate dissolved in concentrated sulphuric acid.

to remove resinous substances, colouring matters, etc., and then digest with alcohol (sp. gr. '85). The solution thus obtained will contain the glucoside and tannin; the latter substance may be removed by precipitation with hide powder or with gelatine. The filtrate will contain the salicin, which on evaporation and cooling will be deposited in the form of crystals.

Salicin is hydrolysed by emulsin to glucose and the alcohol saligenin according to the following equation:—

$$C_{13}H_{18}O_7 + H_2O = C_6H_{12}O_6 + C_6H_4OH CH_2OH$$

Saligenin Saligenin

By the action of sulphuric acid and potassium bichromate salicin is oxidized to salicylic aldehyde C_0H_4OHCHO ; this substance is a fragrant colourless liquid, b.p. 196°, which occurs in the essential oil of *Spiræa Ulmaria*; it is soluble in water, the solution giving an intense violet coloration with ferric chloride; salicylic aldehyde stains the skin yellow.

By employing dilute nitric acid as the oxidizing agent, salicin is converted into helicin, a glucoside which on hydrolysis yields glucose and salicylic aldehyde:—

$$\begin{array}{cccc} C_{13}H_{18}O_7 \,+\, O \,=\, C_{12}H_{16}O_7 \,+\, H_2O \\ Salicin & Helicin \\ \\ C_{12}H_{16}O_7 \,+\, H_2O \,=\, C_6H_{12}O_6 \,+\, C_6H_4OHCHO \\ Helicin & Salicylic aldehyde \end{array}$$

The investigations of Weevers tend to show that ordinarily in the decomposition of salicin, saligenin is really an intermediate substance, the ultimate products being glucose and Thus salicase splits salicin into glucose and saligenin, saligenase produces catechol from the saligenin, and when the leaves decay a third enzyme, catecholase, produces from the catechol an amorphous black pigment. He found that in places where depletion of salicin was taking place the saligenin appeared in quantities insufficient in amount to account for the whole of the salicin; also that catechol occurred in such places after the glucoside had disappeared, in a sufficiently large quantity to warrant the above conclusion. Weevers considers that when glucose and catechol are produced the sugar is translocated, whilst the catechol remains in situ, and combines with fresh glucose, and so reconstructs salicin.

INDICANE.

Indicane,* $C_7H_6NC \cdot O \cdot C_6H_{11}O_5$, is the name given to a glucoside which occurs not only in *Indigofera anil, I. arrecta, I. tinctoria*, and *I. sumatrana*, but also in other plants, such as *Isatis tinctoria*, *Polygonum tinctorium*, species of *Phajus* and other orchids, e.g. *Calanthe* and *Strobilanthes*. In the plant, indicane is well distributed in the aerial organs. Thus in *Indigofera*, it is found in all the tissues of the leaf except the tracheæ of the xylem, it is also abundant in the apex of the stem in all tissues except the wood vessels and the laticiferous system. The flowers also have a small quantity, but the root is characterized by its absence,†

At one time it was considered that the chloroplasts played an important direct part in the formation of indicane, but Leake can find no evidence of this.

Identification.

- 1. The tissue may be boiled in a 2 per cent solution of ammonia. The addition of chloroform to the filtered extract may be made to separate the indigo; the chloroform will rise to the top of the solution, carrying with it the indigo.
- 2. Tissues containing indicane on exposure to the vapour of alcohol for twenty-four hours will turn blue; the reaction will be better marked if the chlorophyll be subsequently dissolved out with absolute alcohol.
- 3. The tissue, in bulk or in section, may be boiled in strong hydrochloric acid and ferric chloride added. The indigo will separate out.
- 4. The tissue is cut up into pieces and quickly immersed in the following mixture:—
 - * The name indicane is also applied to a compound of the formula

This substance, which is more correctly described as indoxyl potassium sulphate, occurs in small quantities in human urine and also in the urine of herbivora.

† Leake: "Ann. Bot.," 1905, 19, 297.

Glacial acetic acid 2 c.c.
Strong sulphuric acid r c.c.
Ammonium persulphate 5 gram.
Water to 100 c.c.

As this fluid penetrates the cells, the indigo is precipitated in blue granules. When penetration is effected fully, the material is washed for twenty-four hours in water, after which sections may be cut and stained in the usual way.

Until a few years ago, *Indigofera* was the only source of the blue colouring matter indigo, for the obtaining of which large tracts of country were under cultivation in India. Within recent years, however, the artificial synthesis of this substance has been effected in a variety of ways and the days of natural indigo are numbered, unless the planters can produce it at a cheaper rate than the chemists.

Indicane, if boiled with mineral acids or hydrolysed by means of an enzyme contained in the plant, breaks up into glucose and indoxyl.

$$C_7H_6NC \cdot O \cdot C_6H_{11}O_5 + H_2O = C_6H_{12}O_6 + C_6H_4$$

$$NH$$
Indicane
$$NH$$
Indoxyl

The latter substance on exposure to atmospheric oxygen undergoes oxidation with the formation of the deep blue colouring matter indigo.

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SECTION IV.

TANNINS.

THE term Tannin is variously employed by different writers, sometimes to denote a particular substance better described as gallotannic or digallic acid, and sometimes as a collective term for a whole group of substances having certain characteristics in common. In order to prevent confusion it is proposed here to use the word tannin only in the latter sense.

The properties of the tannins may be summarized as follows:—

- 1. They are mostly uncrystallizable colloidal substances with astringent properties.
- 2. They precipitate gelatine from solution and form insoluble compounds with gelatine yielding tissues, a property which enables them to convert hide into leather.*
- 3. They all give blackish-blue or blackish-green colours with ferric salts, a fact which is made use of in the manufacture of ink.
- 4. They are precipitated from solution by many metallic salts such as copper or lead acetates or stannous chloride, etc.
- 5. They are precipitated from solution by a strong aqueous solution of potassium bichromate or by a 1 per cent solution of chromic acid.
- *According to some authors this property is not an essential characteristic of tannins; on the other hand Dekker prefers to regard those substances which do not give this reaction as pseudo-tannins and includes under this heading caffetannic acid and the tannins of Portlandia grandiflora, Asperula odorata, Rubia tinctorum, Scrophularia nodosa, Humulus Luphulus, etc. Similarly Procter points out that such substances as moringatannic acid, or maclurin, and caffetannic and lupulotannic acids, are more closely related to the colouring matters than to the tannins; maclurin which is a pentahydroxybenzophenone is the yellow colouring matter occurring in the substance known as fustic, obtained from the wood of Morus tinctoria (see formula on p. 203).

193

- 6. They precipitate from solution both alkaloids and substances of a basic nature, such as basic organic colouring matters.
- 7. In alkaline solution the tannins, and many of their derivatives, readily absorb oxygen, becoming dark in colour.
- 8 With a solution of potassium ferricyanide and ammonia they give a deep red colour.

It must be borne in mind, however, that none of these reactions, taken separately, are specific for tannins; they may be given by many other substances as well, but all true tannins answer them as a whole.

OCCURRENCE.

Tannin, using the word as a generic term, is generally looked upon as an aplastic substance, and is very widely distributed in the vegetable kingdom.

In certain Algæ, e.g. Spirogyra, Mesocarpus and Zygnema, it occurs in the cells in the form of numerous small vesicles; in the Fungi, tannin is stated to be more abundant in parasites than in saprophytes, thus hardly any occurs in the Agaricineæ whilst in the Polyporeæ it is present in much larger amounts.

In the higher plants it occurs more or less generally throughout a tissue, for example in bark, or it may be restricted, in the more mature parts, to special cells which may be isolated or superposed one above the other in the form of chains.

Amongst the higher plants there is no great phylum in which tannin is not found; it occurs in the ferns, e.g. *Angiopteris* and *Aspidium*; in Gymnosperms, e.g. *Pinus*; and also in innumerable Angiosperms, in all parts.

Thus it obtains in the roots of Trianea, Desmanthus and Pistia; in the stems, where it may be accumulated, especially in the bark, of Quercus, many species of Cæsalpinia, Eucalyptus occidentalis, Castanea and Humulus; in the leaves of Cerasus, Rhus, Ficus, and Rhododendron; in the fruit, especially if unripe, of Terminalia Chebula, Cæsalpinia coriaria, Pyrus, and Phaseolus; and more rarely in the seeds, either before or after germination, of Areca Catechu, Echium vulgare and other Boraginaceæ.

Further, tannin is often found in more or less special structures, e.g. the cells of the pulvini of *Mimosa pudica* and *Robinia pseudacacia*; in the gland cells of *Sarracenia* and *Utricularia*; in the hairs of *Primula* and *Hedera*; and also in laticiferous tissue.

Finally, it may be remarked that it is especially abundant in pathological growths such as galls, which may contain from 25 to 75 per cent of tannin.

Kraemer* has investigated the galls formed by the agency of *Cynips aciculata*, a gall fly, upon *Quercus coccinea*. He found that during the chrysalis stage gallic acid was produced, probably at the expense of the starch, and as the imago developed the gallic acid gave place to tannic acid.

In the cell, the tannin occurs in solution in the cell sap, and since tannin forms a precipitate with albuminous matter it follows that the layer of protoplasm around the tannin vesicles must be impermeable to it; if this were not so the protoplasm would be tanned on the production of tannin.

Economically tannin is of great value although it is perhaps not so extensively used at the present day as in the past. As is well known its principal use is in the making of leather †; in this connexion salts of chromic acid have come into use as a substitute, especially in the manufacture of the cheaper grades of leather. Formerly tannins were almost exclusively used in the manufacture of black ink, whereas at the present time various preparations of aniline dyes are in vogue. Among other uses for tannins may be mentioned their value for medicinal purposes and their use as mordants in certain dyeing operations.

The chief sources of tannins are the bark or the wood of various species of Acacia, Castanea, Eucalyptus, and Quercus; the bark of the mangrove Rhizophora Mangle, the roots of Rumex hymenosepalus (Canaigre) and the leaves of Rhus Coriaria (Sumach).

Gallotannic acid is obtained from galls, especially the galls which occur on *Quercus lusitanica*; but the galls on other species of oak, e.g. *Q. sessiliflora*, *Q. pedunculata*, *Q. Ilex*, *Q.*

^{*} Kraemer: "Bot. Gaz.," 1900, 30, 274.

[†] An excellent account of leather manufacture will be found in Procter's "The Making of Leather," Cambridge, 1914.

Cerris, Q. Coccinea, etc., and other plants, e.g. species of Tamarix, are used to a greater or lesser extent.

The amount of tannin present in certain plants varies according to the physiological state, the season of the year, and the conditions of growth.

In *Pinus* it is stated that the amount of tannin varies with that of the resin; thus in the spring it was found that as the tannin decreased in amount so the resin increased. Peacock* found that in *Heuchera americana* the tannin was most abundant in October and least in May, whilst the amount of starch present was greatest in March. Trimble and Peacock found that in *Geranium maculatum* the maximum amount of tannin obtained in April, i.e. just before the period of flowering. From this phase onwards there was a gradual decrease until the minimum was reached in October.

It is found that the more vigorous trees yield the most tannin, and that the character of the soil appears to be of importance. It has been found that oak trees grown in a poor dry soil yield a bark richer in tannin than those grown on the soil of damp lowlands.

According to the observations of Henri, a calcareous soil is more beneficial with regard to tannin formation than is a siliceous soil.

It is not impossible that the different yields of tannin given by the same plant grown in different situations may be due to the relative abundance of the mineral food-materials; thus it has been found that in some instances, e.g. in *Spirogyra* and *Phaseolus multiflorus*, the formation of tannin is inhibited by the absence of chlorine.

With regard to seasonal variation in the amount of tannin in the bark of the oak, the following estimations are given by Eitner:—†

				Q. pedunculati	a. Q. sessiliflora.
April				14.8 per cen	t 12.86 per cent
May				10.41 "	10.46 ,,
June				12.33 ,,	10.58 ,,
July				9.8 ,,	8.11 "
August	t			11.53	10.24

^{*} Peacock: "Amer. Journ. Pharm.," 1891, 172. † Eitner: "Der Gerber, Vienna," 1878, 4.

For the inner bark of the American oak, *Quercus Prinus*, Trimble * found the following seasonal variation:—

December			9.33 per cent
March .			10.63 " "
June .			11.55 " "
July .			11.70 ,, ,,
September			6.66

As a general rule the barks collected in May and June are the richest in tannin, but this does not hold for all parts of plants. Thus, Levi and Wilmer † found that in the case of the horse-chestnut, Aesculus Hippocastanum, the youngest leaves were richest in tannin, the minimum amount obtained in June, whilst in August the quantity rapidly rose until the original value was reached; finally a diminution of tannin occurred just before leaf-fall. Weekly analyses of leaves were made from the opening of the buds to the fall of the leaves in September. The obtained percentages of tannin were: 6.5, 3.3, 3.5, 2.8, 3.7, 3.2, 1.9, 2.8, 3.5, 3.6, 3.4, 5.1, 3.1, 5.3, 4.4, 4.3, 3.4, 6.2, 6.6, 5.2, 6.1, 6.5, 4.5 per cent.

These variations in the tannin-content of parts of plants are of great interest; the value, however, of such estimations would be greatly enhanced if estimations were carried out at the same time to see whether, for instance, there is any obvious relationship between the tannin-content of leaves and of other parts of the plants such as the periderm.

MICROCHEMICAL REACTIONS OF TANNINS.

Before passing on to the detailed examination of the various tannins, the following microchemical tests may be mentioned, but it must be borne in mind that these reactions do not enable one to distinguish between the various tannins.

- 1. Tannins reduce Fehling's solution.
- 2. They are precipitated by basic lead acetate and the salts of many other metals; thus uranium acetate gives a brown precipitate or a brown or brown-red coloration, and an aqueous solution of copper acetate gives a brown precipitate.
 - 3. Potassium bichromate in a strong aqueous solution or

^{*} Trimble: "The Tannins," Philadelphia, 1892, 1894. † Levi and Wilmer: "Hide and Leather," 1905.

a I per cent solution of chromic acid gives brownish-coloured precipitates.

- 4. A red-brown to brown coloration is obtained by the use of a dilute ammoniacal solution of potassium ferricyanide. This test is very delicate, and the reagent must be used sparingly since the coloration is destroyed by an excess.
- 5. The addition of a neutral solution of ferric chloride gives a blue black or greenish coloration or precipitate. Moeller recommends the use of a solution of anhydrous ferric chloride in anhydrous ether.
- 6. A solution of ammonium molybdate in a strong solution of ammonium chloride gives a copious yellow precipitate with many tannins; when added to digallic acid a red coloration results. According to Gardiner* this reagent affords a means of distinguishing glucoside tannin from tannic acid.

The red yellow colour obtained by adding ammonium molybdate to tannic acid is destroyed by oxalic acid.

- 7. Lime water gives a white precipitate which turns red, brown or blue.
- 8. Aqueous solutions of various organic bases such as caffeine and antipyrin precipitate the tannins.

Van Wisselingh† recommends I per cent aqueous solutions of antipyrine and of caffeine.

It must be remembered that several other substances besides tannins are precipitated by these reagents.

- 9. Pfeffer has drawn attention to the fact that tannins are precipitated by methylene blue without prejudice to the vitality of the cells. The stain must be used in very dilute solutions (1 pt. in 500,000 of water), and the tissue under investigation must remain in a large quantity of the solution for several hours. Van Wisselingh's experience is contrary to Pfeffer's, for ne finds that even very dilute solutions of methylene blue are harmful to *Spiregyra*, the plant used by Pfeffer, and after treatment for several days only a little of the tannin was precipitated.
- 10. On the addition of a solution of gelatine a dirty white precipitate is formed.

^{*} Gardiner: "Proc. Camb. Phil. Soc.," 1883, 4, 387. + Van Wisselingh: "Konin. Akad. v. Wetensch., Amsterdam," 1010, 685.

- 11. A brilliant red colour, even when the tannins are in a very dilute solution, results from the addition of an aqueous solution of iodine in potassium iodide mixed with a little 10 per cent ammonia.
- 12. According to Moore,* the action of Nessler's solution (a saturated solution of mercuric iodide in a solution of potassium iodide and potash) varies.
 - (a) A brown precipitate is formed immediately, e.g. epidermis of primrose leaf.
 - (b) A yellow colour is produced which turns reddish brown; finally a brown precipitate comes down, e.g. stem of Yew and Aucuba japonica.
 - (c) A yellow coloration. The compound produced readily diffuses through the cell wall, e.g. young stem and hairs of the ivy.

The following are microchemical tests for gallic acid:

- I. The rapidity of the reaction with potassium chromate may provide a means of distinguishing gallic acid from tannic acid, for in the case of the former a precipitate immediately comes down, whilst in the case of tannic acid, according to Drabble and Nierenstein, the reaction is either very slow or entirely negative.
- 2. Potassium cyanide in aqueous solution gives a pink coloration with gallic acid.
- 3. With Nessler's solution gallic acid gives a grey-green precipitate.

With this same reagent pyrogallol immediately yields a brown precipitate; pyrocatechol forms a deep green precipitate which changes to greenish brown; and a dirty green precipitate is given by protocatechuic acid.

Vinson† recommends exposing the material to be examined to the vapour of nitrous ethers in order to fix and stain the tannin in vegetable tissues. There is, ordinarily, no necessity to cut up the tissue, and as the tannin is deposited in the cells in which it occurs, the method is very convenient for tracing the distribution of the substances in question. The resulting colour varies; thus, the juice of unripe grapes gives a dense brown precipitate; the juice of the persimmon a wine-

^{*} Moore: "Journ. Linn. Soc., Lond., Bot.," 1891, 27, 527. † Vinson: "Bot. Gaz.," 1910, 49, 222.

red coloration; tannic and gallic acids yield a yellow tint, phloroglucin red, and so on.

The material to be examined is merely exposed to the vapour of ordinary sweet spirits of nitre which contain 4 per cent of the ethyl nitrate, or a 20 per cent alcoholic solution of the commercial nitrous ether may be employed. If the latter method be used the time required for full precipitation is considerably less.

CHEMISTRY.

We have as yet comparatively little certain knowledge concerning the chemical constitution of even the simplest tannins. Thus, for example, although ordinary tannic or gallotannic acid is generally regarded as an anhydride formed by the removal of one molecule of water between two molecules of gallic acid, according to the equation

$${}_2C_7H_6O_5$$
 - H_2O = $C_{14}H_{10}O_9$
Gallic acid Gallotannic acid

there are even now, as will be seen below (p. 215), differences of opinion with regard to the exact position from which the two atoms of hydrogen and one of oxygen have been removed.

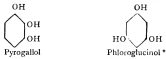
While the composition of the various classes of tannins of course varies considerably, they are probably all more or less complex derivatives of gallic or ellagic acids, or their methylated derivatives, or are condensation products of these or similar acids, with various phenolic substances.

In view of these facts the classification and properties of the tannins will be more easily understood if preceded by a brief description of certain relatively simple phenolic substances from which the complex tannins are built up (p. 220). The substances include the following:—

1. The Dihydric phenols—pyrocatechol, resorcinol and hydroquinone.

2. The dihydroxy acid—Protocatechuic acid.

3. The Trihydric phenols-Pyrogallol and phloroglucinol.



4. The trihydroxy acid—Gallic acid.

The above substances occur in varying proportions among the products obtained by subjecting different tannins to fusion with caustic potash or other chemical treatment; and upon their occurrence is based the chemical classification of the tannins.

PYROCATECHOL, CATECHOL OR PYROCATECHIN. C₆H₄(OH)₂.

This substance is so called from the fact that it is obtained by the destructive distillation of catechu, the sap of *Mimosa Catechu;* it is also obtained by the fusion with potash of other tanno-resins such as kino, the sap of various species of *Pterocarpus, Butea* or *Eucalyptus;* also it occurs in small quantities combined with sulphuric acid in the urine of horses and of human beings. It crystallizes from benzene in colourless glistening plates and melts at 140°.

*Although behaving normally as a trihydric phenol, phloroglucinol reacts with certain ketonic reagents as though it had the following constitution:—

Reactions

- 1. Pyrocatechol is precipitated from aqueous solution by lead acetate. (Distinction from resorcin and hydroquinone.)
- 2. With ferric chloride it gives a green colour which is changed to violet on the addition of sodium acetate.
- 3. Like pyrogallol it reduces silver nitrate in the cold and has therefore been used as a photographic developer.
 - 4. It reduces Fehling's solution on warming.

RESORCINOL. C₆H₄(OH)₂.

This is isomeric with pyrocatechol (for constitutional formula see page 200); it does not generally occur in tannins* but in certain resins, notably galbanum resin and asafætida.

It is used commercially in the manufacture of dye-stuffs, and when heated with sodium nitrite gives the indicator known as Lacmoid.

Resorcinol crystallizes from benzene in colourless needles and melts at 119°; it is somewhat soluble in water, the solution having a sweetish taste.

Reactions

- 1. It is not precipitated from solution by lead acetate.
- 2. With ferric chloride it gives a dark violet colour which is destroyed by the addition of sodium acetate.
- 3. It reduces ammoniacal silver nitrate or Fehling's solution on warming.

HYDROQUINONE.

This third isomer of the formula $C_0H_4(OH)_2$ likewise is not found in tannins, but occurs combined with glucose in the glucoside arbutin and uncombined in the leaves and flowers of *Vaccinium Vitis Idæa*. Hydroquinone crystallizes from water in colourless prisms and melts at 169-170°.

Reactions.

- 1. It gives no precipitate with lead acetate.
- 2. Ferric chloride gives no colour but oxidizes it to quinone.
- * According to Nierenstein, it is produced together with protocatechuic acid and phloroglucinol from quebracho tannin by potash fusion.

- 3. It reduces ammoniacal silver nitrate and Fehling's solution.
- 4. It turns brown in alkaline solution when exposed to the air; its powerful reducing properties enable it to be used in photography as a developer.

PROTOCATECHUIC ACID.

Protocatechuic acid is closely related to pyrocatechol, differing from this substance only by one atom of carbon and two of oxygen which it loses when heated above its melting point (199°), thus:—

$$C_6H_3(OH)_2COOH = C_6H_4(OH)_2 + CO_2$$

Protocatechuic acid Pyrocatechol

It rarely occurs uncombined except, for example, in the fruits of *Illicium religiosum*; in combination, it is found in such substances as Catechin and Maclurin,* both of which give protocatechuic acid on potash fusion; it may further be obtained by a similar process from many resins such as gum benzoin, asafætida, myrrh and also from kino.

Finally its dimethyl ether, known as veratric acid, $C_6H_3(OCH_3)_2COOH$, occurs together with the alkaloid veratrine in the seeds of *Veratrum sabadilla*.

Protocatechuic acid is soluble in water and melts at 199°.

Reactions.

- Aqueous solutions of protocatechuic acid are precipitated by lead acetate.
- 2. It gives a green colour with ferric chloride; on addition of very dilute sodium carbonate the green colour changes first to blue and then to red.
- 3. Ferrous salts produce with protocate chuic acid a violet colour. $\,$
- *Maclurin, sometimes also called moringatannic acid, is a colouring matter of fustic; its constitution is represented by the formula

PYROGALLOL OR PYROGALLIC ACID. C6H3(OH)3.

This substance is so called because it is formed by heating gallic acid according to the reaction—

 $C_6H_2(OH)_3COOH = C_6H_3(OH)_3 + CO_3$ Gallic acid Pyrogallol

It is also formed by fusing hæmatoxylin with caustic potash.

Pyrogallol crystallizes in colourless needles or plates melting at 132° and is soluble in water; its solution, in excess of caustic alkali, absorbs oxygen with great avidity, turning brown and forming carbon dioxide, acetic acid and other substances.

Pyrogallol reduces salts of silver, mercury, or gold to their respective metals.

Reactions.

- 1. Pyrogallol is precipitated from solution by lead acetate but not by lead nitrate.
- 2. It gives a blue colour with ferrous sulphate and a red colour with ferric chloride.
- 3. Aqueous or alcoholic solutions of pyrogallol, in common with those of gallic acid or tannic acid, are coloured purple by iodine.
- 4. Lime water added to an aqueous solution of pyrogallol produces a purple colour which rapidly becomes brown.
 - 5. Solutions of pyrogallol give no precipitate with gelatine.
- 6. Potassium cyanide gives a reddish-brown coloration, which turns brown, but the red tint becomes apparent again on shaking.

PHLOROGLUCINOL. C6H3(OH)3.

Phloroglucinol, which is isomeric with pyrogallol, is produced by fusing a number of resins, such as catechin, kino, dragon's blood, etc., with potash, and it occurs naturally, in a number of glucosides, such as phloretin, quercetin, hesperidin, etc. It crystallizes with two molecules of water, but loses them if heated to 100°, and melts at 218°; it dissolves readily in water, forming a sweet solution, and is readily soluble in alcohol or ether.

Reactions.

- 1. Phloroglucinol is precipitated from solution by lead acetate.
 - 2. It gives with ferric chloride a bluish-violet colour.
- 3. A solution of phloroglucinol in hydrochloric acid produces a red colour on a pine wood shaving; this reaction can also be made use of for detecting lignified cell walls (p. 145).
 - 4. It is a reducing agent, and reduces Fehling's solution.

In addition to the above-mentioned phenols, which are products of the decomposition of tannins by heat or by fusion with alkalis, there are other important substances produced by acid hydrolysis, namely, gallic and ellagic acids and the phlobaphenes.

GALLIC ACID.

Gallic acid, $C_6H_2(OH)_3COOH$, was first prepared by Scheele in 1786 by leaving an aqueous extract of gall nuts to stand in a warm place, and from time to time removing the layer of mould which formed on it; the crystalline precipitate which deposited from the solution was purified by recrystallization from water. Gallic acid, besides occurring in gall nuts, both free and in the form of its anhydride tannic acid, is also found free in sumach, divi-divi, the fruits of $Cesalpinia\ coriuria$, in the leaves of $Arctostaphylos\ Uva-ursi$, and in tea and wine. It may be prepared from tannic acid by acid hydrolysis.

Gallic acid crystallizes in silken needles, and melts at 220°, forming pyrogallol and evolving carbon dioxide; it is sparingly soluble in cold water, but dissolves readily in hot water and in alkalis; alkaline solutions, like those of pyrogallol, absorb oxygen from the air, becoming brown in colour; they also reduce metallic solutions of silver or gold and Fehling's solution.

Gallic acid is converted into its anhydride digallic acid by heating with phosphorus oxychloride to 130° or by boiling with arsenic acid:—

$$\begin{array}{lllll} 2C_6H_2(OH)_3COOH & = & C_{14}H_{10}O_9 & + & H_2O \\ & & & \text{Digallic acid} & \end{array}$$

This digallic acid precipitates gelatine from solution, and for

this reason it was regarded by Schiff* as being identical with gallotannic acid. This view has been proved by Walden† to be untenable, since the physical properties of the two substances are quite different (see p. 212).

Reactions.

- 1. Gallic acid is precipitated from solution by lead acetate; on adding caustic potash a carmine-coloured precipitate is formed, which dissolves in excess to a raspberry-red solution.
- 2. Ferric chloride produces a blue-black colour or precipitate according to the strength of the solution; excess of ferric salt changes the colour to green, while excess of gallic acid reduces the ferric salt to ferrous and destroys the colour.
 - 3. Iodine solution produces a transient red colour.
- 4. Gallic acid does not precipitate gelatine from solution. (Distinction from tannic acid.)
- 5. When heated with concentrated sulphuric acid it turns green and then purple, being converted into rufigallic acid, $C_{14}H_sO_8$, a substance used in dyeing.
- 6. Potassium cyanide gives a pink colour which disappears on standing, but returns again on shaking with air.
- 7. Lime water gives a blue coloration or precipitate; in very dilute solutions a reddish colour is produced.

ELLAGIC ACID. C14H6O8.

Closely related to gallic acid is the substance known as Ellagic acid, its name being obtained by the inversion of the word gallic.

Whether or not this substance occurs free in nature is not definitely established; certain it is, however, that ellagic acid can be readily obtained by the hydrolysis of ellagitannic acid, a substance which almost invariably accompanies gallotannic acid in the numerous vegetable products in which this latter occurs; it also occurs in conjunction with tannins of the pyrogallol class, and constitutes the bloom which is produced on leather by this type of tannin.

^{*} Schiff: "Ber. deut. chem. Gesells.," 1871, 4, 232, 967; 1879, 12, 33; "Annalen," 1873, 170, 143. + Walden: "Ber. deut. chem. Gesells.," 1897, 30, 3153; 1898, 31, 3167.

The most convenient natural sources are "divi-divi" (Cæsalpinia coriaria), "algarobilla" (Cæsalpinia brevifolia), "myrobalans" (Terminalia Chebula), etc. Aqueous extracts of these fruits on long standing frequently deposit ellagic acid, most probably by the action of a ferment contained in the plant; it is, however, prepared by pouring a hot concentrated alcoholic extract of divi-divi into cold water; the acid is thereby precipitated, and may be filtered and purified.

Synthetically it may be prepared by the oxidation of gallic acid by means of arsenic acid, or better by oxidizing gallic acid in acetic acid solution with potassium persulphate and sulphuric acid.*

Ellagic acid is a yellow microcrystalline solid which is very slightly soluble in water, and therefore readily separates from aqueous solutions in which it is formed; it is also very slightly soluble in alcohol or ether, but dissolves somewhat readily in boiling pyridine. The dried substance treated with 1-2 drops of nitric acid gives, on dilution with 10-20 drops of water, a blood-red colour (Griessmayer's reaction).

Its constitution is, according to Graebe, best represented by the formula—

from which it will be seen that it may be considered to be produced by the abstraction of two molecules of water from two molecules of gallic acid with simultaneous oxidation or removal of two atoms of hydrogen.

Ellagic acid is used to some extent as a dye-stuff, being sold under the name of "Alizarine yellow in paste," for use with chromium mordants.

Catellagic, Metellagic, and Flavellagic acids are the names given by Perkin to artificially synthesized acids obtained by him. They are closely related to ellagic acid, but have not, as yet, been found to occur naturally.

^{*} Perkin and Nierenstein: "J. Chem. Soc., Lond.," 1905, 87, 1415.

PHLOBAPHENES.

Among the products of the decomposition of tannins by boiling with acids must be mentioned the substances known as Phlobaphenes. The name derived from the Greek ($\phi \lambda \omega i \delta s$ —bark, and $\beta a \phi \hat{\eta}$ —dyeing) was first given by Stahelen and Hoffstetter,* in 1844, to a red-brown substance obtained by them by adding water to an alcoholic extract of bark which had previously been extracted with ether to remove fats or waxes. It has since been shown that aqueous extracts of bark containing tannin, deposit from solution a substance known as oakred or phlobaphene, and that this substance is more rapidly produced by warming concentrated solutions of tannin with sulphuric acid.

Inasmuch as phlobaphenes are produced by any process which tends to remove water, such as heating tannins to a high temperature or prolonged boiling or heating under pressure, they are regarded as anhydrides of the tannins; besides being thus produced artificially, they occur also in nature side by side with the tannins from which they can be produced.

They are red-coloured substances and are practically insoluble in water though they dissolve in solutions containing tannic acid; also they dissolve in alcohol and in alkaline solutions.

The formation of phlobaphenes by treatment of a tannin with acid is characteristic of pyrocatechol tannins (p. 211) in just the same way as ellagic acid is produced from pyrogallol tannins.

A number of different phlobaphenes are known, such as kino-red, catechu-red, oak-bark red, etc.

TANNINS AS GLUCOSIDES.

At one time it was thought that the tannins were substances of a glucosidic nature and occurred in the plant in combination with a carbohydrate complex such as glucose; but while this is undoubtedly so in some cases it is by no means universal.

To determine whether a tannin is a glucoside or not the following procedure is recommended by Procter.†

The Tannin must first be carefully purified from glucose

* Stahelen and Hoffstetter: "Annalen," 1844, 51, 63.

Procter: "Leather Industries Laboratory Book, London," 2nd ed., 1908.

gums, or other bodies likely to interfere. This may be done by extraction according to Pelouze's method (p. 213), or, if the tannin is to be extracted from an aqueous solution, by agitating with ether to remove gallic acid and then saturating the aqueous solution with common salt and shaking with ethyl acetate, which extracts the tannin. The ethyl acetate is then evaporated off, the last traces being expelled by the repeated addition of small quantities of ether.

Another method is to extract with alcohol and to evaporate off the alcohol at as low a temperature as possible, and then to take up the residue with a large volume of water whereby the phlobaphenes are precipitated and may be filtered off. The infusion is then precipitated with successive small quantities of lead acetate; the first and last portions are rejected and the middle fraction after washing is suspended in water and saturated with sulphuretted hydrogen. The precipitated lead sulphide is filtered off, and the solution is warmed to drive off excess of gas and then extracted with ethyl acetate.

Thus purified the tannin or its washed lead salt is heated to 100° for an hour or more in a sealed tube or boiled in a flask under a reflux condenser with hydrochloric acid (2 per cent). After cooling, the mixture is allowed to stand for some time and is then filtered from any deposit which may have formed. The filtrate is shaken with ether to remove gallic acid and the aqueous solution boiled, neutralized with caustic soda and precipitated with basic lead acetate to remove any unchanged tannin or colouring matter; the solution is again filtered and any lead remaining in solution is removed by the addition of dilute sulphuric acid, excess of acid being carefully avoided. The solution is then neutralized and once more filtered and the clear filtrate heated to boiling with Fehling's solution when a red precipitate proves the presence of glucose.

THE CLASSIFICATION OF TANNINS.

With the present incomplete state of our knowledge concerning the chemical constitution of the tannins, it is difficult to make a proper chemical classification of these substances.

According to Trimble * the tannins may be divided into two main groups:—

^{*} Trimble: loc. cit., vol. ii. p. 132.

I. Those containing about 52.2 per cent carbon.

This group includes the tannins contained in oak galls (gallotannic acid), chestnut (wood and bark), sumach, pomegranate, etc.

2. Those containing about 59-60 per cent carbon.

This group includes taunins of oak bark, kino, canaigre, ratanhia, catechu.

He points out that the fact of similar percentage composition would not itself be sufficient to justify this classification, but he finds that the classification still holds when the reactions towards certain reagents are compared as under:—

	Group 1.	Group 2.		
Ferric salts.	Blue colour and precipi- tate.	Green colour and pre- cipitate.		
Lime water.	White precipitate be- coming blue.	Light pink precipitate becoming red and brown.		
Bromine water.	No precipitate.	Yellow precipitate be- coming brown,		

Dekker * proposes the following classification:-

- 1. Catechin tannins, occurring in gambier, catechu and *Hamamelis* bark.
 - 2. True tannins-
 - (a) Gallic acid group . . . gallotannic acid; tea and sumach tannins.
 - (b) Ellagic acid group . . . divi-divi, algarobilla and myrobalan tannins.
 - (c) Oak bark group . . . the majority of red-producing tannins.
- 3. Pseudotannins (which do not form leather with hide), caffetannic acid and the tannins of maté, hops, etc.

Perhaps the best classification is the one given by Procter,† who divides tannins into two main groups:—

(A) Pyrogallol tannins, including divi-divi, galls, sumach, myrobalans, valonia, algarobilla, oak gall, oak wood and chestnut tannins.

These tannins have the following characteristics:-

- 1. They give with ferric salts a dark blue colour.
- 2. They give no precipitate with bromine water.
- They produce on leather a "bloom" consisting of ellagic acid.
- (B) Pyrocatechol tannins, including all the pine barks,
- * Dekker: "De Looistoffen," Amsterdam, 1906.
- + Procter: "The Principles of Leather Manufacture," London, 1903.

acacias, mimosas, oak barks (but not oak wood, fruits or galls), quebracho wood, cassia and mangrove barks, canaigre, cutch and gambier.

The tannins of this class are characterized by the following properties:— $\dot{}$

- 1. They give with iron alum a greenish-black colour, though the reaction is liable to be rendered uncertain by the presence of other colouring matters.
- 2. When treated with bromine water, until the solution smells strongly of it, they give a yellowish or brown precipitate; in weak solutions the precipitate may form slowly.
- 3. The addition of concentrated sulphuric acid to a drop of the infusion produces a dark red or crimson ring at the junction of the two liquids; on dilution the liquid turns pink.
- 4. These tannins deposit no "bloom," but when boiled with acids deposit red insoluble colouring matters known as phlobaphenes (see p. 208).

Some of the tannins belonging to this group, notably gambier and cutch, contain phloroglucinol as one of their constituents; this substance may be tested for by moistening a pine wood shaving with a little of the infusion and then adding a little concentrated hydrochloric acid; the formation, after a short time, of a bright red or purple stain indicates the presence of phloroglucinol.

PROPERTIES AND DESCRIPTION OF INDIVIDUAL TANNINS.

As already stated the term Tannin is merely a generic name for the whole group of substances, though it has been, and still is, frequently used to mean a particular tannin, namely that contained in oak galls. This substance is, however, better named gallotannic acid, as it is customary to name the tannins after the source from which they are obtained; thus quercitannic acid indicates the tannin of oak bark, sumactannin that derived from sumac, and so on.

PYROGALLOL TANNINS.

GALLOTANNIC ACID.

(Syn. Digallic acid, or Tannic acid, or merely "Tannin".)

The two chief commercial sources of gallotannic acid are :-

- 1. Turkish or Aleppo galls, produced by the gall wasp Cynips gallæ, which lays its eggs in the buds of Quercus infectoria. These contain from 50 to 60 per cent of gallotannic acid.
- ¹ 2. Chinese galls, produced by the burrowing of *Aphis chinensis* in the leaf-stalks of young twigs of *Rhus semialata*. These galls may contain up to 70 per cent of gallotannic acid.

Gallotannic also occurs in sumach (*Rhus Coriaria*), in tea and in many other plants.

Until recently there was some difference of opinion as to whether this tannin occurs in the plant combined with glucose in the form of a glucoside, or whether the sugar which is frequently found associated with it is merely an impurity.*

E. Fischer and Freudenberg,† on reinvestigating the question, have found that "tannin," even after repeated careful purification, yielded about 7 to 8 per cent of glucose on hydrolysis with sulphuric acid; from this it is concluded that "tannin" or gallotannic acid is in reality a compound of five molecules of digallic acid with one molecule of glucose, in which the five hydroxyl groups of the sugar are esterified by five molecules of acid. Such a compound would be a pentadigalloyl glucose of the formula—

$$C_6H_7O_6[C_6H_2(OH)_3CO.O.C_6H_2(OH)_2.CO]_5 \ or \ C_{16}H_{52}O_{46}$$

The authors have, moreover, been able to synthesize a substance having a similar structure by esterifying glucose by means of tricarbomethoxy-galloyl chloride—

$$\begin{split} &C_6H_{12}O_6+5C_6H_2(OCOOCH_3)_3COCI=C_6H_7O_6[C_6H_2(OCOOCH_3)_3CO]_5+5HCI \\ \text{and saponifying the resulting compound with caustic soda:} &-- \end{split}$$

$$C_6H_7O_6[C_6H_2(OCOOCH_3)_3CO]_5 \\ \Longrightarrow C_6H_7O_6[C_6H_2(OH)_3CO]_5$$

This pentagalloyl glucose closely resembles "tannin" as regards optical activity, feebly acid character, solubility, taste, colour with iron salts and ability to precipitate gelatine or alkaloids. This furnishes a striking confirmation of the correctness of Fischer's suggested constitution of "tannin," and it

†Fischer and Freudenberg: "Ber. deut. chem. Gesells.," 1912, 45, 915 and 2709.

^{*}Cf. Strecker: "Annalen," 1852, 81, 248; 1854, 90, 328; Pottevin: "Compt. rend.," 1901, 132, 704.

is to be hoped that the synthesis * of "tannin" itself may soon be effected.

EXTRACTION OF GALLOTANNIC ACID.

Gallotannic acid is best prepared by extracting finely-powdered gall nuts with a mixture of twelve parts of ether with three parts of alcohol; twelve parts of water are then added and, after shaking, the lower aqueous layer is run off from below and evaporated. The resulting tannic acid may be decolorized by boiling with animal charcoal.

Pelouze recommends the following method: The powdered material is heated under a reflux condenser with a mixture of thirty parts of ether, five parts of water, and two parts of alcohol. On cooling three layers of liquid are formed, of which the lowest contains 33 per cent, the middle 8 per cent, and the top 2 per cent of the tannic acid present in the substance.

Gallotannic acid forms an amorphous powder† which, when pure, is almost colourless; it is readily soluble in water, forming a solution with an astringent taste and which reacts acid to litmus; it dissolves also in alcohol or glycerine, but is only sparingly soluble in ether and is insoluble in chloroform, benzene, ligroin or carbon disulphide; it is also insoluble in hydrochloric or sulphuric acids and is precipitated by these substances from its aqueous solutions; it is soluble in alkalis, and the solution, as in the case of gallic acid or of pyrogallol, rapidly absorbs oxygen from the air and darkens in colour.

When boiled with 2 per cent hydrochloric acid for some time, gallotannic acid is broken up into gallic acid.

If heated slowly from 160 to 215° and kept at the higher temperature for thirty minutes, carbon dioxide, water, pyrogallol and metagallic acid are produced. The pyrogallol volatilizes and condenses in the cooler part of the vessel.

The action of heat on tannins may also be studied by dissolving I gram of tannin in 5 c.c. of glycerine, heating slowly to 210° and maintaining the liquid at this temperature for half

^{*}See also Fischer and Strauss: id., 1912, 45, 2467.

[†]What is known as "Crystal tannin" in commerce is not really crystalline; it is made by drawing a syrupy solution into threads and breaking these up after drying.

an hour. The liquid is then cooled and shaken with 20 c.c. of ether; after the addition of water the ethereal solution is separated and evaporated; the residue contains pyrogallol.

Reactions.

- Ferrous sulphate, free from ferric salts, produces at first no change, but on exposure to air the solution darkens from oxidation.
- 2. Ferric chloride produces a blue-black colour or precipitate.
- 3. A dilute solution of iodine in potassium iodide gives a transient pink colour, as in the case of gallic acid.
- 4. Gallotannic acid is precipitated from solution by gelatine, and similarly combines with hide powder converting it into leather. (Distinction from gallic acid.)
- 5. Gallotannic acid precipitates proteins, alkaloids and many other organic substances from solution.
- \acute{o} . Lead nitrate or lead acetate gives precipitates of lead tannate. (N.B.—Neither pyrogallol nor gallic acid is precipitated by lead nitrate, though both give precipitates with lead acetate.)
- 7. Potassium cyanide gives a reddish-brown colour which changes to brown, but the red tint reappears on shaking with air.
 - 8. Lime water gives a grey precipitate.

DETECTION OF GALLIC ACID IN PRESENCE OF GALLOTANNIC ACID.

Gallic acid may be detected in the presence of gallotannic acid by dissolving the mixture in water and extracting the solution with ether; the ethereal extract on evaporation yields gallic acid which may be identified by the usual tests.

Gallotannic acid may also be separated from gallic acid by adding a solution of lead acetate strongly acidified with acetic acid; under these circumstances lead tannate is precipitated while lead gallate remains dissolved.

Similarly tannic acid is precipitated by many alkaloids and basic substances which have no action on gallic acid.

THE CONSTITUTION OF GALLOTANNIC ACID.

The close relationship subsisting between gallotannic and gallic acids was first observed by Scheele, who, by allowing an

infusion of gall nuts to undergo fermentation, obtained gallic acid. Within recent years this change has been studied anew by Fernbach,* who isolated a tannin splitting enzyme, tannase, from Penicillium, and also by Pottevin, + who isolated a similar enzyme from the mould Aspergillus.

This change, which may be represented by the equation— $C_{14}H_{10}O_9 + H_2O = 2C_7H_6O_5$

may be effected more rapidly by boiling gallotannic acid with dilute sulphuric acid.

When, therefore, it was found by Schiff that gallic acid could be converted back into the anhydride by means of phosphorus oxychloride it was assumed that this substance, which was called digallic acid, was identical with "tannin". This view came to be generally accepted, although objections were raised from time to time on the ground that the physical constants, such as electrical conductivity and absorption of light, of natural tannin and synthetic digallic acid were different.§

The formula which Schiff in 1871 assigned to his synthetic product was as follows:-

but it was not until Dekker | pointed out that natural tannic acid was optically active, that real doubt was cast on this formula. Dekker accordingly proposed an alternative formula containing an asymmetric carbon atom as follows:-

Other suggestions were offered by Iljin ¶ and by Nierenstein **

- * Fernbach: "Compt. rend.," 1900, 131, 1214. † Pottevin: "Compt. rend.," 1900, 131, 1215.
- \$Schiff: "Ber. deut. chem. Gesells.," 1871, 4, 232.

 Walden: "Ber. deut. chem. Gesells.," 1897, 30, 3151; 1898, 31, 3167.

 Dekker: "Ber. deut. chem. Gesells.," 1906, 39, 2497.
- ¶ Iljin: "Journ. Prakt. Chem.," 1900, (2), 82, 422.
- ** Nierenstein: "Ber. deut. chem. Gesells.," 1908, 41, 77; 1909, 42, 1122; 1910, 43, 628.

to account for the discrepancies above mentioned, but in view of E. Fischer and Freudenberg's work, already referred to above, these suggestions are valueless.

Incidentally it may be mentioned that two other digallic acids are known in addition to Schiff's acid, namely:—

- 1. β -digallic acid, $C_{14}H_{10}O_9 + 2H_2O$, obtained by heating ethyl gallate with pyroracemic acid, and
- 2. Fischer's * digallic acid, $C_{14}H_{10}O_9$ or $C_{14}H_{10}O_9 + C_7H_6O_5$, obtained by the action of tricarbomethoxygalloylchloride with dicarbomethoxygallic acid.

ELLAGITANNIC ACID.

This tannin, which is commonly found together with gallotannic acid, is important as being the mother substance of ellagic acid, which is responsible for the bloom characteristic of pyrogallol tannins. The quantity of this substance present in different plants varies considerably; it is greatest in dividivi. Amongst the other tannins giving ellagic acid bloom may be mentioned algarobilla, myrobalans, chestnut tannin, pomegranate tannin, valonia, etc.

Ellagitannic acid, unlike ellagic acid (p. 207), is soluble in water or alcohol; prolonged boiling with water converts it into ellagic acid. It has been variously described by different authors as a glucoside, as a hydrated soluble form of ellagic acid, or as a condensation product of ellagic acid with gallic acid,†

PYROCATECHOL TANNINS.

CATECHU TANNIC ACID.

Catechu tannic acid is the name given to the tannin contained in gambier catechu and in Bombay catechu or cutch, a substance obtained by evaporating an aqueous extract of the bark of various trees (see below). A similar tannin is also contained in kino.

Little is known as to its constitution, but it is believed to be an anhydride of catechin.

^{*}Fischer: "Ber. deut. chem. Gesells.," 1908, 41, 2890; "Annalen," 1911,

⁺ Cf. Nierenstein: "Ber. deut. chem. Gesells.," 1907, 40, 4575; 1909, 42, 353; 1910, 43, 1257.

[#]This substance is used largely for dyeing.

CATECHIN.

This substance, which is obtained from Acacia catechu, Ouroparia catechu, mahogany wood, Mimosa, and pyrocatechol tannins in general, is not in itself a tannin since it does not precipitate gelatine, but it is converted into a tannin, namely catechu tannic acid, by loss of water, a change which may be rapidly brought about by heating to 120° or above.

Catechin may be prepared by extracting powdered catechu with ether; the crude material obtained on evaporating off the ether may be purified by crystallization from water.

Catechin forms colourless glistening needles, which, when dry, melt at 175-177°. It is readily soluble in alcohol and ethyl acetate, not so readily soluble in ether, and only slightly soluble in cold water.

With ferric chloride alone it gives a green colour, but with ferric chloride and sodium acetate a dark violet.

It gives the phloroglucin reaction with pine wood shaving and hydrochloric acid.

Potash fusion gives protocatechuic acid and phloroglucinol.

QUERCITANNIC ACID.

Quercitannic acid is the name given to the tannin of oak bark, which is not identical with the tannin of oak galls.

Pure quercitannic acid yields no glucose on hydrolysis, though levulose is nearly always present in oak bark.

Although much work has been done on the oak bark tannins by various workers, notably Etti, Böttinger and Löwe, nothing definite is known as yet regarding their constitution.

Procter summarizes the present state of our knowledge by saying that, on the whole, it seems probable that the principal tannin of oak bark is a purely catechol tannin, and that the gallic and ellagic acids which have been detected in it are due to an admixture of the gallotannic and ellagitannic acids present in oak wood.

A great many more tannins are known, but too little is known about their composition to justify their inclusion here.

PHYSIOLOGICAL SIGNIFICANCE OF TANNINS.

It is manifestly a difficult matter to ascertain the significance of tannins in the life of the plant, more especially as these substances vary in different species, so that what may be true for one is not necessarily true for all.

It is, therefore, not surprising to find that several ideas have been put forward.

With regard to the origin of tannins practically nothing of fundamental importance is known.

According to the investigations of Kraus, tannin, although not a direct photosynthetic product—as is indicated by the fact that the tannin does not increase in the leaves of plants which are able to photosynthesize in dull light—is not formed unless carbon dioxide and light are available. He found that etiolated leaves produced no tannin, and that the amount of this substance in shaded leaves was less than that contained in the leaves of the same plant fully exposed to the sun. The tannin thus formed is translocated to the stem and root.

This, however, is not the only origin for tannin, for if tannin-containing seeds, e.g. the oak, be germinated in darkness, there is an increase in the amount of tannin; further, the production of various aromatic compounds may be a stage in the synthesis of proteins, and some of these may eventually give rise to tannin.

The facts regarding the distribution of tannin have an important bearing on the subject. It is abundant in leaves; in parts in which growth is very active, such as growing points; in galls and other pathological growths; also it is found in association with secretory organs, such as gland cells of Sarracenia and Utricularia, and in parts in which the protoplasm is especially irritable, such as pulvini. Pfeffer found that in young fully formed pulvini no tannin occurs, but it appears soon after movements commence and gradually increases in quantity until the leaf dies.

In the case of *Robinia pseudacacia* the pulvini of the leaflets contain less tannin than the main pulvinus, which is much less sensitive than are the secondary pulvini.

The consideration of these facts supports the conclusion arrived at by Sachs that tannin results from intense metabolism such as occurs in active leaves; in rapid tissue formation, as in galls and vegetative apices; during germination and secretion; and as a consequence of particular stimulation, as in mobile pulvini.

Various facts on the relation between tannin and other substances such as starch, sugar, resin, etc., have led to various opinions.

That starch frequently is contained in the same cells with tannin suggests a connexion between the two, and it is not impossible that the starch may contribute the glucose for the construction of the tannin. In the case of *Pinus*, it has already been mentioned that in the spring, when the flow of resin is most copious, the tannin decreases as the resin increases; also the cells surrounding the epithelium of resin ducts contain tannin and starch. Wiesner, therefore, concluded that tannin is an intermediate product in resin formation.

Tannin is not uncommon in unripe fruits, and the amount of these astringent substances diminishes during ripening.

According to Bassett* "the amount of tannin in fruits varies with certain factors, such as injury, length of time between removal from tree and analysis, etc. The presence and relative amount of this tannin or tannin-like body is controlled by the presence of certain enzymes which vary in amount and activity during the growth of these fruits."

Buignet, from the fact of the diminution of tannin and starch which occurs concurrently with the increase in sugar, considered that the sugar in the ripe fruit (e.g. Musa) is formed from these two substances. This opinion, however, is not held by Gerber who investigated the same phenomenon. In Diospyros Kaki he found the young fruit to be very astringent, but not so the ripe fruit. He considers that the tannins disappear by complete oxidation without the formation of carbohydrates. One reason for his opinion is that in the conversion of tannin into carbohydrate more carbon dioxide would have to be liberated than oxygen absorbed, whereas in fruits the relation is the reverse.

On the other hand, he does consider the tannins to be of some value, for they, by the formation of pectins, may limit the loss of carbohydrate.

Further, inasmuch as the pleasing odours of fruit are ac-

^{*}Quoted from the footnote appended to a paper on the Toxicity of Tannin by Cook and Taubenhaus: "Delaware Coll. Agric. Exp. Station" Bull., 91, 1911.

quired after the tannin has disappeared it is not impossible that the latter may have some connexion with the formation of fruit esters.

Other suggestions regarding the value of tannin are not wanting; thus Moore* states that it may play an important part in the lignification of cell walls.

More recently Drabble and Nierenstein† have come to the conclusion that tannins play an important part in cork formation, and are acted upon in the plant by formaldehyde and acids and are precipitated in the walls of the cork cells. Reasons for this theory may be alluded to briefly.

There occur in plant tissues tannins; phenols such as phloroglucinol, resorcinol and hydroquinone; and hydroxybenzoic acids, such as gallic, salicylic, and protocatechuic acids.

When these substances are treated with hydrochloric acid and formaldehyde various condensation products are precipitated. These condensation products can be produced from gallic acid, pyrogallol, protocatechuic acid, phloroglucinol, salicin, tannic acid, and other substances, simply by passing a slow stream of carbon dioxide through the mixture of formal-dehyde and the tannic acid, for example.

The reactions given by these bodies are similar to those characteristic of cork; thus they are insoluble in Schweitzer's reagent and strong sulphuric acid, but readily dissolve in strong potash. It is, therefore, possible that in cork formation similar condensation products may play a part, for the requisite materials are present in the plant.

Further, in the plants examined, the presence of gallic or tannic acids was indicated in the immediate neighbourhood where cork was being formed, and by suitable means there can be obtained from cork, products having the same mother substance as the condensation products mentioned above.

Still more recently Van Wisselingh has published certain observations from which he concludes that tannin plays an important part in the formation of cell walls in certain cases, for instance *Spirogyra*. He does not consider it a reserve foodmaterial as such, but rather a soluble substance which the

^{*} Moore: loc. cit.

[†] Drabble and Nierenstein: "Biochem. Journ.," 1906, 2, 96.

t Loc. cit.

plant makes use of in elaborating other materials. This conclusion is in agreement with the opinions held by Wingand and published in 1862. Van Wisselingh worked with *Spirogyra*, and the main facts on which he based his conclusions are as follows. Cells which are about to conjugate are rich in tannin, and as the process of conjugation proceeds, there is a gradual diminution in the amount of this substance, so that the mature zygospore contains nothing more than mere traces.

If conjugation be interrupted at an early stage, there is still an increase of tannin, so that when death results there is relatively a large quantity present. This accumulation may be used as an argument in support of the view that tannin is a waste product. Van Wisselingh, however, remarks that this should not be a source of wonder, for in this case "It is not the intention of Nature that it should be wasted. Nature ensures a sufficient supply of tannin in *Spirogyra*, because this substance is required in development, as for instance in conjugation and spore-formation. The occasional failure to conjugate as a result of which much tannin is lost, does not prove that it is a waste product and not a plastic material."

The author in question also found that a diminution of tannin occurred during the formation of the cell wall after nuclear division, and if the tannin were precipitated during the earliest phases of cell division, the cell wall was not formed although the nucleus divided into two quite normally. Cladophora, which does not contain tannin, was used as a control; it was found that by keeping the filament in a solution of antipyrine, the reagent used in the experiment on Spirogyra, the cell-wall formation was not disturbed.

It must be mentioned that Van Wisselingh does not claim that tannin is the only substance used in cell-wall formation, nor does he maintain that the only physiological significance of tannin is its use as a plastic material.

Finally, in this particular connexion, it may be mentioned that tannin may play a part in the formation of various pigments such as anthocyan and erythrophyll, for similar decomposition products (compounds allied to the phenols) may be obtained from each.

The fact that some Fungi can make use of tannin as a food material provided that it is not in excess, and the facts that

many are glucosides, and that oxidation readily takes place with the ultimate formation of oxalic acid and carbon dioxide, suggest that the substances under consideration may be reserve food-material.

Thus Schell, while acknowledging that tannin may sometimes be a bye-product of metabolism, considered that at other times it might be used up in the construction of higher compounds which would serve as food. He found that, in the germination of the oil-containing seeds of *Echium vulgare* and other Boraginaceae, as the oil is used up the tannin begins to play a part in the constructive metabolism and gradually diminishes in amount. Further, if such seeds be germinated in the light the tannin increases in quantity. For these and other reasons he concluded that such a use of tannin only obtained when there was a scarcity of the more normal foods such as starch and oil.

A consideration, however, of other facts does not tend to support the idea of tannin being of the nature of a reserve Hillhouse,* for example, found that if a fuchsia having an abundant supply of tannin be grown in the dark, there is no diminution in the substance in question. Then again the facts of its distribution are against this particular view; for example, it does not occur in sieve tubes which transport both sugar and other food substances; there is, in many cases, not a great discrepancy in the tannin-content of fully mature and fallen leaves, for naturally it would be expected that if tannin were of any considerable value as a food-stuff it would not be accumulated in bark and old leaves but would be translocated out of such places before they were cast off, the same as are other materials in the generality of cases. But against this argument may be cited the fact that fallen leaves may contain substances of undoubted value to the plant, such as nitrogen and phosphorus, and even glucose and starch. In evergreen leaves there is no diminution in the quantity of tannin during the winter months, which may mean that either it is of no great value or that, since growth is more or less at a standstill, the plant has more food than it requires immediately, or that it subserves some biological function; thus Warming has suggested that in

^{*} Hillhouse: "Midland Naturalist," 1887-8.

this particular connexion the tannin may be of value in protecting the plant against undue evaporation during the winter, and further it may be a means of rapidly restoring lost turgor.

On the other hand the figures obtained by Levi and Wilmer, mentioned above, require some explanation; why should a minimum of tannin occur in the leaves in June when photosynthesis is so very active? is it used up in the construction of other substances or is it merely translocated to other parts such as the bark? If the latter be true, the further question arises, then why should it be transferred at one time of the year and not at another?

Of course, it is possible that these and like variations may be explained by the varying conditions of, say, light, temperature and moisture; and with regard to this variation in the amount of tannin, more especially in germinating seeds, Van Wisselingh points out that the amount found at any particular moment represents the balance as it were of the tannin account; that is to say, if more tannin is formed than is decomposed, an increase in the tannin content will result and vice versa, so that in one and the same plant there will be sometimes an increase and sometimes a decrease according to the conditions obtaining. It does not necessarily follow, and this is applicable to many things besides tannin, that because there is an increase in the amount, therefore the substance is of no value in constructive metabolism.

Tannin has been considered an important constituent of the osmotic substances of the cell; although this may be true for some tannins it probably does not hold for all, since no ill effects follow the precipitation of tannins in the living cell by means of methylene blue; also, in certain cases, it is not renewed when precipitated.

A biological significance is not infrequently attached to tannins; thus it may be of use against animals, it may be connected with the activity of nectaries in providing sugar, and it has been suggested by Moore that when it occurs in the epidermis of leaves, it may play a part in the opening and closing of stomata.

Finally, it may be of considerable value as an antiseptic, preventing the germination and growth of parasitic Fungi. In

this connexion Cook and Taubenhaus* have found that in many cases tannin has a tendency to retard or inhibit the growth of Fungi, the parasitic forms being more sensitive than the saprophytic. In some cases the spores are killed, whilst in others germination is much impeded. On the other hand, low percentages of tannin may in some instances stimulate germination and also fruiting. The behaviour of Fungi towards tannin varies with the species and sometimes even with the individual, more especially in the case of spores.

To conclude, the different substances included under the term Tannin are so numerous as to make it improbable that they all have the same physiological significance.

[&]quot;Cook and Taubenhaus: "Delaware Coll. Agric. Exper. Station" Bull., 91, 1911.

SECTION V.

PIGMENTS.

CHLOROPHYLL.

As is well known, chlorophyll is contained in the chloroplasts which are universally present in green plants and vary considerably in their size, shape, and number within the cell. With regard to their structure there has been much dispute. It is, however, generally agreed that the structure of the plastids is either reticulate or vacuolate.

The pigment itself is variously stated to be dissolved in some oily substance which is held in the channels and meshes of the plastids, or to exist in the form of a precipitate; and with regard to the distribution of the pigment within the plastid there is again some dispute. According to many, it is distributed evenly throughout the stroma, whilst, on the other hand, others maintain that it is restricted to the peripheral layers of the plastid.

Amongst the most recent contributions to the subject is the investigation of Priestly and Irving* on the chloroplasts of certain species of *Selaginella* and *Chlorophytum*. They find that the pigment is restricted to the peripheral regions of the chloroplast, where it is held in the meshes of the network of the matrix. They agree with Timiriazeff's views that the function of the chlorophyll necessitates its distribution in very thin layers in order that the amount of energy set free may be as great as possible.

With regard to the origin of the chloroplast there is also some dispute. The general view, due originally to Schimper and Meyer, appears to be that plastids do not arise *de novo* within the cell, but by the division of pre-existing plastids, so that, in this respect, there is continuity between parent and offspring. This has led to the conception that originally the

^{*} Priestly and Irving: "Ann. Bot.," 1907, 21, 407.

chloroplasts once had a separate individuality, and that, in a sense, ordinary plants are parasitic upon the imprisoned plastids which have become permanent members of the structures of the cell.

On the other hand, other investigators hold that the chloroplasts may arise from differentiated parts of the protoplasm, which parts are not plastids. Lewitski* draws attention to the presence of minute bodies occurring in the protoplasm, but not in the nucleus, which he calls mitochondria, chondriosomes, etc. These structures, which he considers are essential parts of the cytoplasm, increase by division, and give origin to the plastids. For instance in the pea, Pisum sativum, and the asparagus, Asparagus officinale, the mitochondria of the cells of the stem apex give rise to chloroplasts, whilst those of the apex of the root are converted into leucoplasts. Meyer,† however, is opposed to these conclusions. finds that very minute chloroplasts occur in the cotyledons of the seed of Helianthus annuus, and increase in size and divide by fission as germination proceeds and maturity is reached.

In green plants chlorophyll may occur not only wherever light gains access to the living cells, but also in places where light seemingly cannot penetrate, at any rate in any quantity, for instance in the cortex internal to the periderm-not only in small twigs, but also of larger branches—in the medullary rays and even in the pith. \ Also it may occur in the cotyledons of seeds before they are set free from the ovary or from the cone; Pinus, Euonymus europæus, and species of Cucurbita are familiar examples. In some of these cases light no doubt does penetrate through the walls of the superposed cells; this may be well seen if the seeds be removed and the lumen of the fruit of the vegetable marrow be cleaned out. It is hardly necessary to remark that if the chlorophyll in these deeply-seated tissues be functional, its contributions to the

^{*}Lewitski: "Ber. deut. bot. Gesells.," 1910, 28, 538. † Meyer: id., 1911, 29, 158. See also Schmidt: "Prog. Rei. Bot.," 1912, 4, 163; Forenbacher: "Ber. deut. bot. Gesells.," 1911, 29, 648; Woycicki, "Sitz. Warschauer Ges. Wiss.," 1912, 5, 167; and Löwschin: "Ber deut.

bot. Gesells," 1913, 31, 203.

† Miller: "Ann. Bot.," 1910, 24, 693. An excellent résumé of the literature on mitochondria is given by Cavers in "New Phyt.," 1914, 13, 96, 170. § See Scott: "Ann. Bot.," 1907, 21, 437.

food-stuffs of the plant, as Goldflus* has pointed out, must be of considerable value.

But in some cases the pigments of such chloroplasts may not be the same as those of the ordinary chloroplasts of the leaf; thus, according to Monteverde and Lubinenko,† the seeds of many Cucurbitaceæ contain not chlorophyll, as ordinarily understood, but chlorophyllogen, which may pass over into chlorophyll under the influence of light and some other factor, possibly enzymic.

Also it must be remembered that it does not follow that because chlorophyll is present, photosynthesis necessarily takes place, even though the requisite conditions, light and supply of raw material, obtain. Thus it appears probable that the chlorophyll in green parasites is not functional, and the same holds for the chlorophyll in the gynacium of certain plants, e.g. *Ornithogalum arabicum*. At any rate, in these cases the photosynthetic power is so small as to be masked by the respiratory activity.

Attention may here be drawn to the work of d'Arbamont, ‡ who considers that the plastids containing chlorophyll may be divided into two classes, chloroplasts and pseudochloroplasts. Of these the former include those bodies usually termed chloroplasts, and are characterized by the fact that they do not swell in water, and do not, as a rule, stain when treated with acid aniline blue. On the other hand, pseudochloroplasts swell in water and do stain with aniline blue. In some cases plants may contain pseudochloroplasts only.§

With regard to the conditions necessary for the formation of chlorophyll, light is the most important, but in addition a certain degree of temperature, as well as the presence of certain substances, such as iron and magnesium, are essential. There is, however, some dispute regarding other factors. Palladin || states that chlorophyll formation is an oxidative process, and, as a result of his experiments, finds that etiolated leaves on

^{*} Goldflus: "Rev. Gén. Bot.," 1901, 13, 49.

[†] Monteverde and Lubinenko: "Bull. Jard. Imp. Bot., St. Pétersbourg," 1909, 9, 27.

[‡]D'Arbamont: "Ann. Sci. Nat. Bot.," 1909, 9, 197.

[§] See Belzung: id., 1891, 13, 17; "Journ. Bot.," 1895, 9, 67, 102.

^{||} Palladin: "Ber. deut. bot. Gesells.," 1891, 9, 194, 229; 1902, 20, 224; "Rev. Gén Bot.," 1897, 9, 385.

exposure to daylight will not form chlorophyll unless a supply of carbohydrate is available. If an etiolated leaf does not contain carbohydrate, then greening will take place if the cut leaf be placed in a solution of sugar. Almost any sugar will do, e.g. sucrose, maltose, glucose, fructose, or raffinose; success was also obtained by the use of glycerine. The solution used must be neither too weak nor too strong; a strong solution of sucrose, for instance, will retard the chlorophyll formation because it will depress oxidative processes. On the other hand, Issatchenko * finds that etiolated leaves of certain plants, e.g. those of *Vicia Faba*, when detached from the plant and placed in strong sugar solution, even 50 per cent, will form chlorophyll. He considers that light is the all-important factor.

With regard to the substances which immediately precede chlorophyll, and from which chlorophyll is formed, nothing definite is known.

The chemical study of chlorophyll dates from the year 1819, when Pelletier and Caventou + first applied this name to the green leaf pigment without, however, isolating the substance. Since then, numerous workers have attempted to prepare chlorophyll in a pure condition, but the methods employed in most cases were of too drastic a nature for the substance to escape destruction. Previous to 1911, there was no chemical evidence to show that chlorophyll was not a single chemical individual, although Stokes,‡ Sorby,§ and others had obtained spectroscopic evidence pointing to the existence of more than one substance; confirmatory evidence was subsequently obtained by Tswett. In 1912, however, Willstätter and Isler ¶ definitely showed that chlorophyll as ordinarily obtained, and to which they had originally assigned the formula C₅₅H₇₂O₆N₄Mg, is in reality a mixture of two substances :-

> Chlorophyll a C₅₅H₇₂O₅N₄Mg ** Chlorophyll b C₅₅H₇₀O₅N₄Mg.

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* Issatchenko: "Bull. Jard. Imp. Bot., St. Pétersbourg," 1906, 6, 20.
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and

[†] Pelletier and Caventou: "Ann. Chim. Phys.," 1819, 9, 194.

[‡] Stokes: " Proc. Roy. Soc.," 1864, 13, 144.

[§] Sorby: id., 1872, 21, 442.

^{||} Tswett: "Ber. deut. bot. Gesells.," 1906, 24, 326; 1907, 25, 137; "Ber. deut. chem. Gesells.," 1908, 41, 1352.

[¶] Willstatter and Isler: "Annalen," 1912, 390, 269.

^{**} For the physical characteristics of these two substances see page 234.

229

Accompanying chlorophyll are three yellow or reddishbrown pigments, Carotin, Xanthophyll, and Fucoxanthin (the latter occurring only in brown algae), which are known collectively as the Carotinoids. Owing to the similarity in solubilities between these substances and chlorophyll, their complete separation is a matter of some difficulty; it was first effected by Willstätter and Hug.*

The average proportions in which these various constituents occur in different plants have been determined by Willstätter, and are approximately as follows:—

		In Land Plants.†	Brown Algæ † (Fucus).	Green Algæ† (Ulva).
Chlorophyll a .		.62	.16	.093
" b.		'22	.01	*o66
Carotin		·055	.0312	*014
Xanthophyll .		'093	10305	·036
Fucoxanthin .			.059	

From these figures the following interesting deductions may be made:—

- I. The molecular proportions between chlorophylls and carotinoids are as 3.5 to I \ddagger in terrestrial plants, but only I to I in the case of algæ.
- 2. In the brown algae chlorophyll a predominates, only about 5 per cent of the mixture being chlorophyll b; in terrestrial plants, on the other hand, the proportion is pretty constantly about 3:1.
- 3. In the green algæ there is relatively more of chlorophyll b.

Concerning the physiological significance of these substances it has been suggested by Willstätter \S that since chlorophyll b ($C_{55}H_{70}O_6N_4Mg$) contains more oxygen than chlorophyll a ($C_{55}H_{72}O_5N_4Mg$), the former compound is produced by the action of chlorophyll a upon carbon dioxide

^{*} Willstätter and Hug: "Annalen," 1911. 380, 177.

⁺ These figures are percentages calculated on the dry material.

^{**} With regard to this ratio, it has been stated by Willstätter that it is remarkably constant, and that there is a greater variation between different leaves of the same plant than between corresponding leaves of different plants. This view is, however, contested by Borowska and Marchlewski ("Biochem. Zeitschr.," 1913, 57, 423), who hold that it is entirely dependent on external circumstances, such as soil, stage of growth, etc.

[§] Willstätter: "Untersuchungen über Chlorophyll," p. 237.

during assimilation, and that chlorophyll b is then reconverted into chlorophyll a with evolution of oxygen. On the other hand, the molecular formulæ of carotin ($C_{40}H_{56}$) and xanthophyll ($C_{40}H_{56}O_2$) only differ by two atoms of oxygen, and the close association between the carotinoids and chlorophyll may be explained by assuming that the function of carotin is to reduce chlorophyll b to chlorophyll a, being itself oxidized to xanthophyll, and that the latter compound is reconverted by some enzyme into carotin with evolution of oxygen.

With a view to throwing some light on the mechanism of photosynthesis, Wager* has studied the decomposition of chlorophyll on exposure to oxygen both in the light and in the dark, with the result that he finds that the process is not catalytic. Oxygen is absorbed and aldehydes are formed, and it is suggested that the sugars produced during assimilation are not formed directly from carbon dioxide and water but by the polymerization of aldehydes produced in this way. Warner† has also found that formaldehyde is produced when chlorophyll is exposed to sunlight or electric light in air; since this substance is produced both in the presence and in the absence of carbon dioxide, it would appear that the latter plays no part in the production of formaldehyde by photosynthesis outside the plant, and that the formaldehyde is in reality an oxidation product of the chlorophyll.

Quantitative measurements of the relation between the amount of carbon dioxide assimilated and the weight of chlorophyll concerned have been made by Willstätter and Stoll. ‡ A regular stream of air containing a known amount of carbon dioxide was passed over from 5 to 20 grams of leaves contained in a small illuminated glass vessel immersed in a constant temperature water-bath. By estimating the amount of carbon dioxide in the issuing gas and the amount of chlorophyll in the leaves, they determined the so-called assimilation number for different leaves which was the ratio between the amount of carbon dioxide assimilated per hour and the weight of chlorophyll concerned in the assimilation. Experiments with normal, autumnal, and etiolated leaves showed that the assimilation is

^{*} Wager: "Proc. Roy. Soc.," 1914, [B], 87, 386.

⁺ Warner: id., 378.

[‡] Willstätter and Stoll: "Ber. deut, chem. Gesells.," 1915, 48, 1540.

not always proportional to the chlorophyll content, which may be explained by assuming that some enzyme takes part in the process. The fact that in leaves rich in chlorophyll increased illumination produces no increased assimilation, whereas a rise in temperature does, is attributed to the accelerating effect of increased temperature upon enzyme action. In the case of leaves deficient in chlorophyll, on the other hand, increase of temperature has but little effect, whereas such leaves are very susceptible to increased illumination. The explanation in this case is that there is more than sufficient enzyme for the chlorophyll, but that the greatest assimilative effect can only be attained when all the chlorophyll is exerting its maximum activity. Attempts to bring about assimilation with chlorophyll outside the leaf failed, presumably owing to the absence of this enzyme. The removal of epidermis from the under surface of leaves had no deterrent effect on assimilation, but a slight pressure applied to the leaves brought assimilation to a complete standstill.

THE CONSTITUTION OF CHLOROPHYLL.

As already stated, chlorophyll was first isolated from its accompanying yellow pigments, the carotinoids, by Willstätter and Hug in 1911, and in the following year it was shown by Willstätter and Isler that the chlorophyll so obtained was not a single substance, but a mixture of two distinct substances, chlorophyll a and chlorophyll b, in the proportion roughly of three molecules of the former to one of the latter. The constitutions provisionally assigned to these two substances are given by the following formula:—

$$\begin{array}{ccc} \text{COOCH}_3 & \text{COOCH}_3 \\ \text{COOC}_{29} \text{H}_{29} & \text{COOC}_{29} \text{H}_{39} \\ \text{COO}_{39} \text{H}_{29} & \text{COOC}_{29} \text{H}_{39} \\ \text{Chlorophyll } a & \text{Chlorophyll } b \end{array}$$

from which it may be seen that they are both esters of methyl and phytyl alcohol ($C_{20}H_{30}OH$), and that the former contains what is known as a lactan grouping.

The recognition of magnesium as an essential constituent of chlorophyll, which is due to Willstätter,* has proved of immense value in the study of the degradation products of chlorophyll.

By the action of alkalies and acids respectively upon the two chlorophylls, it has been found possible to divide the degradation products of chlorophyll into two groups:—

- I. Those that retain magnesium, known as Phyllins.
- 2. Those that are free from magnesium, known as Porphyrins.

The Action of Alkalies.

When the two chlorophylls are treated with the calculated amount of concentrated methyl alcoholic potash their ester groups are hydrolysed, and two isomeric tribasic acids result from each which are known as chlorophyllin and isochlorophyllin a or b, as the case may be:—

$$\begin{array}{c|cccc} \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{COOCH}_{3}}{\mathbf{COOC}_{20}\mathbf{H}_{29}} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg}}{\mathbf{COOH}} & \overset{\mathbf{COOH}_{3}}{\mathbf{COOH}} \\ & & & & & & & & \\ \mathbf{Chlorophylli} \ a & & & & & & \\ \mathbf{Chlorophyllin} \ a & & & & & & \\ \mathbf{C}_{32}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{COOH}_{3}}{\mathbf{COOH}} \\ & & & & & & & \\ \mathbf{C}_{32}\mathbf{H}_{29}\mathbf{O}_{2}\mathbf{N}_{4}\mathbf{Mg} & \overset{\mathbf{COOH}_{3}}{\mathbf{COOH}} \\ & & & & & & \\ \mathbf{C}_{32}\mathbf{H}_{29}\mathbf{O}_{2}\mathbf{N}_{4}\mathbf{Mg} & \overset{\mathbf{COOH}_{3}}{\mathbf{COOH}} \\ & & & & & \\ \mathbf{C}_{32}\mathbf{H}_{29}\mathbf{O}_{2}\mathbf{N}_{4}\mathbf{Mg} & \overset{\mathbf{COOH}_{3}}{\mathbf{COOH}} \\ & & & & & \\ \mathbf{C}_{32}\mathbf{H}_{29}\mathbf{O}_{2}\mathbf{N}_{4}\mathbf{Mg} & \overset{\mathbf{COOH}_{3}}{\mathbf{COOH}} \\ & & & & & \\ \mathbf{C}_{32}\mathbf{H}_{29}\mathbf{O}_{2}\mathbf{N}_{4}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} \\ & & & & \\ \mathbf{C}_{32}\mathbf{H}_{29}\mathbf{O}_{2}\mathbf{N}_{4}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} \\ & & & \\ \mathbf{C}_{32}\mathbf{H}_{29}\mathbf{O}_{2}\mathbf{N}_{4}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} \\ & & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} \\ & & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} \\ & & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} \\ & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{N}_{3}\mathbf{Mg} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} \\ & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{H}_{29} \\ & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} \\ & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{H}_{29} \\ & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} \\ & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{H}_{29} \\ & & \\ \mathbf{C}_{31}\mathbf{H}_{29}\mathbf{Ng} & \overset{\mathbf{C}_{31}\mathbf{H}_{29}\mathbf{H}_{29} \\ & & \\ \mathbf{C}_{31}\mathbf{H}_{31}\mathbf{H}$$

Chlorophyllin a when heated with alkali loses carbondioxide, and yields two isomeric dibasic acids, glaucophyllin and rhodophyllin, $C_{31}H_{32}N_4Mg(COOH)_2$, and at a higher temperature it loses two molecules of carbon dioxide, yielding a monocarboxylic acid, pyrrophyllin, $C_{31}H_{33}N_4Mg(COOH)$. By heating with soda lime the third molecule of carbon dioxide may be removed with the formation of aetiophyllin a substance containing no carboxyl group at all, and to which the following formula is assigned:—

^{*} Willstätter: "Annalen," 1906, 350, 48.

Aetiophyllin C31H34N4Mg

The Action of Acids.

Acids, especially oxalic acid, remove magnesium from all derivatives containing this element, replacing it by two atoms of hydrogen without altering the rest of the molecule.

Thus chlorophyll a and b give by removal of Mg the compounds

Phæophytin
$$a$$
 $C_{32}H_{32}ON_4$ $COOC_{20}H_{39}$ and $COOC_{20}H_{39}$ $COOCH_3$ $COOCH_3$ $COOCH_3$ $COOCH_3$ $COOCH_3$ $COOCH_3$ $COOC_{20}H_{39}$

respectively, while chlorophyllin a gives phytochlorin f and g, $C_{32}H_{32}ON_4(COOH)_2$. On the other hand, glauco and rhodophyllin by removal of magnesium give glauco and rhodoporphyrin $C_{31}H_{31}N_4(COOH)_2$, while pyrrophyllin yields pyrroporphyrin $C_{31}H_{35}N_4(COOH)$. By removing the last carboxyl from the latter compound a substance aetioporphyrin $C_{31}H_{35}N_4$ is obtained, which is the magnesium free analogue of aetiophyllin $C_{31}H_{34}N_4Mg$.

CRYSTALLINE AND AMORPHOUS CHLOROPHYLL.

The physical constants of these substances as determined by Willstätter and his pupils are as follows:—

Chlorophyll (a and b). Analysis agrees with formula C55H72O6N4Mg. Bluish-black glistening powder, with metallic

Appears crystalline under the microscope. No definite M.P.

Practically insoluble in cold light petroleum, but dissolves readily on addition of a few drops of methyl or ethyl alcohol.

Phase Test.

(i.e. hydrolysis in ethereal solution, with methyl alcoholic potash), gives a transient brown coloration (cf. p. 240).

Chlorophyll a. C55H59O5N4Mg.

Bluish-black powder.

Bluish-black powder.

Sinters and forms a viscous mass at 117-121°

Very sparingly soluble in light petroleum, but dissolves very easily in most organic solvents.

Phase Test.

Transient pure yellow Transient colour.

Chlorophyll b. CssHzoO.N.Mg.

Dark green or greenishblack glistening powder.

Dark green or greenishblack glistening powder. Sinters at 86-92°, and becomes viscous at 120 130°.

Quite insoluble in light petroleum, and is generally somewhat soluble than chlorophyll a.

Phase Test.

brilliant red colour.

From the above data it will be seen that neither ordinary chlorophyll (a and b) nor either of the constituents of this mixture show any marked tendency to crystallize which at first sight would appear to be in contradiction with the wellknown fact first observed by Borodin* that when green leaves are moistened with alcohol, and allowed to evaporate slowly under a coverslip, crystals of chlorophyll may be observed under the microscope. Willstätter and Benz† described a method of obtaining this substance in quantity from Galeopsis tetrahit, and later Willstätter and Stoll t showed that this so-called crystalline chlorophyll was not present as such in the plant, but was a secondary product produced by the action of the alcohol upon the chlorophyll under the action of an enzyme chlorophyllase. The phytyl group is thereby replaced by the ethyl group as illustrated by the equation:—

$$\begin{array}{c} \text{COOCH}_3\\ \text{COOC}_{21}\text{H}_{29}\text{N}_3\text{Mg} & \text{COOC}_{20}\text{H}_{39} + \text{C}_2\text{H}_5\text{OH} = \text{C}_{31}\text{H}_{29}\text{N}_3\text{Mg} & \text{COOC}_2\text{H}_5 + \text{C}_{20}\text{H}_{39}\text{OH} \\ \text{NH} & \text{NH} \\ \text{Amortphous Chlorophyll a} & \text{Crystalline Chlorophyll a} \end{array}$$

Crystalline Chlorophyll a

* Borodin: "Bot. Ztg.," 1882, 40, 608.

+ Willstätter and Benz: "Annalen," 1907, 358, 267.

Willstätter and Stoll: id., 1910, 378, 18.

For the monomethyl ester of chlorophyllin a Willstätter has proposed the name chlorophyllide a,

and adopting this nomenclature, amorphous chlorophyll would be termed phytylchlorophyllide, while crystalline chlorophyll would be ethylchlorophyllide.

Chlorophyllase belongs to the same class of enzymes as lipase; the latter substance, however, is only able to hydrolyse amorphous chlorophyll, replacing the phytoxyl group by hydroxyl; it cannot effect alcoholysis.

On the other hand, working with methyl alcohol and chlorophyllase, it has been found possible to replace the phytyl group by methyl, forming methylchlorophyllide, which is the methyl analogue of ethylchlorophyllide or crystalline chlorophyll; it is best obtained by treating fresh leaves with 50-60 per cent methyl alcohol; if prepared from *Hevadeum* it is sparingly soluble in ether and crystallizes from that solvent in steel-blue glistening prisms; that prepared from stinging nettles is slightly less soluble in ether and crystallizes in triangular and hexagonal plates.

By acting in moist ethereal solution in the absence of alcohol, ordinary hydrolysis was effected with the formation of the monomethyl ester of the chlorophyllin, namely, chlorophyllide—

$$\begin{array}{c|c} C_{31}H_{29}N_3Mg & COOCH_3\\ & COOH\\ & CO\\ NH \end{array}$$

this is an extremely unstable substance which forms green plates.

The enzyme is sensitive to high temperatures, and when boiled with alcohol it is gradually destroyed; its activity is greater at 25° than at 30°.

Chlorophyll appears to be always accompanied by the enzyme, the amount increasing with the amount of chlorophyll, and hence young leaves appear to contain less enzyme than the older ones. In *Pyrus Aucuparia*, *Mellitis Melissophyllum*, *Stachys silvatica*, *Lamium maculatum*, and *Heracleum* the

amount of enzyme is comparatively large. *Urtica, Avena,* ordinary grasses, *Sambueus, Platanus, Aspidium, Equisetum,* and *Taxus* may be conveniently employed for demonstrating the effect of the enzyme by leaving the tissues in contact with an alcoholic extract of amorphous chlorophyll; practically all the phytol is thereby removed.

The enzyme is also able to effect the synthesis of phytyl chlorophyllide (amorphous chlorophyll) from chlorophyllide and phytol.

The constitution of this alcohol phytol has been studied by Willstütter, Meyer, and Hüni,* who find it to be unsaturated and have provisionally assigned it the formula

RELATIONSHIP BETWEEN CHLOROPHYLL AND HÆMOGLOBIN

With a view to the further elucidation of the constitution of the chlorophyll molecule, especially in regard to the complex to which the carboxyl groups are attached, the oxidation of the porphyrins by means of chromic acid in the presence of sulphuric acid has been studied by Marchlewski† and by Willstätter and Asahina.‡ These investigations point to the

existence of the grouping $\begin{bmatrix} C-C \\ C-C \end{bmatrix} N$ in the molecule, since

the two chief oxidation products are found to be pyrrole derivatives of the formulæ—

The former substance, which is the imide of a tricarboxylic acid known as hæmatinic acid, of the formula—

$$\mathbf{CH_3.\,C-COOH}$$

$$\mathbf{COOH.\,CH_2.\,CH_2.\,C-COOH}$$

^{*} Willstätter, Meyer, and Hüni: "Annalen," 1910, 387, 73.

[†] Marchlewski: "Chem. Zentralbl.," 1902, I, 1017. ‡ Willstätter and Asahina: "Annalen," 1910, 373, 227.

has also been obtained from hæmoglobin, the red colouring matter of the blood, and a connexion between hæmoglobin and chlorophyll is thereby established.

The relationship between this hæmatinic acid imide and hæmoglobin is as follows:—

Hæmoglobin is readily hydrolysed by dilute acids or alkalis with the formation of hæmatin; this latter substance contains iron, which can, however, be readily removed by treatment with hydrogen bromide in acetic acid solution,* giving an iron free compound hæmatoporphyrin;† both hæmatin; and hæmatoporphyrin on oxidation yield the hæmatinic acid imide mentioned above.

Another link between chlorophyll and hæmoglobin is supplied by the fact that Willstätter and Asahina § have obtained from chlorophyll by reduction three pyrrole derivatives:—

one of which, hæmopyrrole, has also been obtained by the reduction of hæmatoporphyrin.

With regard to the manner in which the magnesium or iron are respectively united to the complex molecules of chlorophyll and hæmoglobin, the following skeletons, involving the assumption of subsidiary valencies, according to Werner and others, have been suggested ||:--

In this connexion compare the formula assigned to Aetiophyllin (page 233).

- * Nencki and Zaleski: "Zeit, physiol. Chem.," 1900, 30, 423.
- †It should be noted that chlorophyll derivatives free from magnesium are by analogy called porphyrins: cf. Phylloporphyrin, etc.
 - # Küster: "Zeit. physiol. Chem.," 1899, 28, 1; 1900, 29, 185.
 - § Willstatter and Asahina: "Annalen," 1911, 385, 188.
 - | Willstätter and Fritzsche: id., 1909, 371, 33.

EXTRACTION OF CHLOROPHYLL.

The usual method of extracting chlorophyll from green tissues consists in first steeping the fresh material in hot water to destroy oxidizing enzymes and then extracting the colouring matter by means of warm alcohol. Willstätter, however, recommends the use of dried in place of fresh material, and extracting by shaking with organic solvents (ethyl or methyl alcohol, ether or acetone) in the cold.

The chief advantage in using dried material lies in the fact of its relatively small bulk, 100 grams of stinging nettle leaves, for example, weighing only 25 grams after drying. It has been shown, moreover, that the operation of drying produces no change of any importance in the chlorophyll, since the results obtained from dried material have been repeated and confirmed on fresh material.

On the other hand, organic solvents containing an appreciable amount of water are preferable to the dry solvents. This is attributed by Willstätter to the fact that aqueous solvents dissolve out salts, such as potassium nitrate, from the cell sap, and these affect the state * of the colloidal solution of chlorophyll in the chloroplast, thereby rendering the chlorophyll more easily accessible to the solvent. Moreover, the number of substances going into solution is thereby increased, and the solution is no longer effected by the solvent alone but by the solvent together with the accessory substances.

If dry solvents are used, the extract is much less pure since it contains a larger proportion of carotinoids, lecithins, etc., whose solubilities are very similar to those of chlorophyll.

The following methods of extracting dried or fresh leaves respectively are described by Willstätter:—

1. Half a kilo of dried material is spread on a porcelain Buchner funnel in a layer of not more than 4 to 5 cms. thick, and 1.5 litres of solvent are drawn through this layer by means of a filter pump in the course of half an hour. This filtrate, measuring about 0.9 litre, contains from 4.25 to 4.5 grams of chlorophyll.

The solvent employed may be either 90 per cent (aqueous) alcohol or 80 per cent (aqueous) acetone. The former filters

^{*} See section on Colloids, page 283.

rather more rapidly, but acetone has the advantage over alcohol in preventing the chlorophyll from undergoing what is known as allomerization, a peculiar change which interferes with its power of crystallization, and prevents it giving the phase test.

2. Two and a half kilos of fresh leaves are ground up in a mill and shaken in a bottle with 1'5 litres of acetone to remove water and mucilage and to stop enzyme action. The acetone is then filtered off on a pump; it contains no chlorophyll. The residue is then freed from acetone by filtering on a pump under a pressure of 200 atmospheres, and the resulting hard mass, weighing 0'8 kg., is broken up and ground again. On adding 1'5 litres of acetone the latter becomes diluted to 80 per cent by the water still remaining in the residue; the mixture is shaken for 5 minutes and a further quantity of 1 litre of 80 per cent acetone is now added. The liquid is filtered off on a pump and the residue treated three times with half a litre of 80 per cent acetone. The total filtrate should measure 3'7 litres and contain 4'7 grams chlorophyll.

In order to ascertain what proportion of the total chlorophyll present has been removed in any particular extraction, another quantity of dried material, say from 100 to 200 grams, may be subjected to an exhaustive percolation with an excess of alcohol until the alcohol comes through colourless. Both extracts are then diluted until 1 kg. of dry powder corresponds to 200 litres of extract and their strengths are compared by means of a colorimeter.

Similarly, a fairly accurate estimate of the amount of chlorophyll present in a solution can be made by colorimetric comparison with a solution containing '025 gram of pure crystallized chlorophyll dissolved in t litre of alcohol. For this purpose the yellow colouring matters must, however, be removed; this is done by allowing the solution to stand for some time with alcoholic potash; the solution is then decanted from the brown resinous deposit which settles on the sides of the vessel, and, after washing the latter with a little more alcohol, the combined alcoholic solutions are diluted with water and extracted with ether to remove the yellow colouring matters.

After suitably diluting with alcohol, the solution is then

compared in a colorimeter with the standard chlorophyll solution.

In this way it was found that I kg. of fresh stinging nettle leaves containing 25.6 per cent of total solid contained an amount of chlorophyll equivalent to 1.6 grams of the crystalline substance, corresponding, therefore, to 1.6 × 1.38 = 2.2 grams of amorphous chlorophyll.*

The following simple experiments are selected from a number described by Willstätter and Stoll† to illustrate the properties of chlorophyll and the carotinoids:—

- I. Grind up 10 grams of fresh stinging nettle leaves with silver sand in a mortar. Cover with 20 c.c. acetone and filter over a pump; wash the residue with more acetone and filter again; the filtrate will contain 0.02 gram chlorophyll.
- 2. Dried powdered leaves do not part with their colour on treatment with benzene or light petroleum, and only yield chlorophyll very slowly to anhydrous alcohol, acetone, or ether, but may be readily extracted by means of 90 per cent alcohol or 80 per cent acetone yielding a green solution with a strong red fluorescence.
- 3. Prepare an ethereal solution of chlorophyll as follows: About 15 c.c. of an 80 per cent acetone extract of dried leaves are poured into 30 c.c. of ether contained in a tap funnel and mixed with 50 c.c. water. The ethereal solution rises to the surface. It should be washed four times with 50 c.c. of water each time by carefully allowing the water to run down the side of the funnel without shaking. If a 30 per cent methyl alcoholic solution of potash is now run under the ether layer a brown colour is produced at the junction of the two liquids. The colour gradually changes to olive-green and finally back to the original green. The reaction, which is known as the "Phase Test," is due to the saponification of the chlorophyll with formation of the potassium salt of chlorophyllin. Consequently on dilution with water the green colour remains in the aqueous layer and is no longer soluble in ether.
- 4. Shake vigorously 5 c.c. of an ethereal solution of chlorophyll (prepared as above) with 2 c.c. of concentrated methyl

^{*}The factor 1'38 for converting crystalline into amorphous chlorophyll represents the ratio between the molecular weights of these two substances.

+ Willstätter and Stoll: "Untersuchungen über Chlorophyll," Berlin, 1913.

alcoholic potash. When the green colour has returned dilute with 10 c.c. water, added in portions, and add a little more ether. On shaking, two layers separate; the lower aqueous alkaline layer contains the chlorophyll while the ether contains carotinoids.

5. To separate xanthophyll from carotin wash the ethereal solution of these two substances obtained from previous experiment with a little water and evaporate to 1 c.c. Dilute with 10 c.c. of light petroleum, and shake up two or three times with 10 c.c. of 90 per cent methyl alcohol until the latter is no longer coloured. The methyl alcohol will contain the xanthophyll while the carotin will be in the light petroleum.

THE CAROTINOIDS OR YELLOW PIGMENTS ACCOM-PANYING CHLOROPHYLL.

In addition to chlorophyll three pigments which are insoluble in the cell sap occur in plants either in a relatively pure form in chromoplasts, or associated with chlorophyll in the chloroplasts; they are carotin, xanthophyll, and fuco-xanthin.

Of these pigments the most important are carotin and xanthophyll, which were at one time supposed to be identical. Thus Tammes* considers that most yellow pigments, whether in separate plastids or associated with the chloroplasts, consist of carotin. From the researches of Arnaud† and Willstätter and Mieg‡ there is no doubt that xanthophyll and carotin are different substances.

Willstätter and Escher, moreover, have isolated from the fruits of the tomato a yellow pigment, lycopin, isomeric with carotin. It differs, however, from carotin in some of its physical properties and in the amount of oxygen it takes up on oxidation.

Although carotin and xanthophyll are commonly associated with chlorophyll they are not antecedents of this substance.

^{*} Tammes: "Flora," 1900, 87, 205.

⁺ Arnaud: "Bull. Soc. Chim.," 1887, 48, 64.

[#]Willstätter and Mieg: "Annalen," 1907, 355, 1.

[§] Willstätter and Escher: "Zeit. physiol. Chem.," 1910, 64, 47.

CAROTIN C40H 56.

This pigment is widely distributed and, as has already been mentioned, is generally associated with chlorophyll in the chloroplasts. It also occurs in various forms, amorphous or crystalline, in various parts of many plants. The colour of yellow or orange petals is not infrequently due to it, e.g. the corona of the common Narcissus, N. Poeticus; similarly the presence of innumerable small intracellular crystals of carotin are responsible for most of the colour of the root of the carrot. and so also is the tint of many fruits where the carotin is often in amorphous granules.

With regard to the physiological significance of carotin, the work of Tammes and Kohl* shows that carotin absorbs certain rays of radiant energy which can be made use of in photosynthesis.

In addition to this there is the possibility that carotin may be of importance in respiration, acting in a manner comparable to the hæmoglobin of the blood.†

The possible function of the carotinoids in assimilation has already been referred to on page 230.

In those cases where a large amount of carotin occurs in organs of storage, such as the roots of the carrot, it may be of value as a reserve food-material. Finally, where the colours of flowers are due to its presence, carotin is important in the floral biology.

Carotin is insoluble in water and very slightly soluble in acetone or cold alcohol; in hot alcohol it is more soluble; and in ether, chloroform, light petroleum, and carbon bisulphide it is readily soluble. The colour of the solution varies from yellow to red; on crystallization flat reddishyellow plates are formed which exhibit the phenomenon of dichroism, being orange-red by transmitted light and greenishblue in reflected light.

According to Willstätter, ‡ carotin may be extracted from stinging nettle leaves by light petroleum; it has the molecular formula C₄₀H₅₆₃ and is probably identical with the sub-

^{*} Kohl: "Ber. deut. bot. Gesells.," 1906, 24, 222. † Arnaud: "Compt. rend.," 1889, 109, 911. ‡ Willstätter and Mieg: "Annalen," 1907, 355, 1.

stances erythrophyll and chrysophyll described by Bougarel and Schunck respectively.

It absorbs 34'3 per cent of its weight of oxygen, being converted into a colourless substance. With iodine it forms the compound $C_{40}H_{56}I_2$, which crystallizes in dark violet prisms.

XANTHOPHYLL C40H56O2.

This substance is closely related to carotin, having the molecular formula $C_{40}H_{56}O_2$. Ewart * has, indeed, shown that xanthophyll may be converted into carotin by the action of zinc dust or magnesium powder and water.

It is a neutral substance, reacting neither as an alcohol nor as an acid.

It absorbs 36.55 per cent of its weight of oxygen, and forms an additive compound with iodine of the formula $C_{40}H_{56}O_2I_2$, which crystallizes in dark violet tufts.

The more important physical constants and solubilities of carotin and xanthophyll are given in the appended table, compiled by Willstätter:—

	-			
			Carotin.	Xanthophyll.
Appearance		•	Copper coloured leaflets.	Pleochroic dark reddish- brown plates,
Colour by	transmi	tted		
light .			Red.	Yellow to orange.
Melting-point			167·5-168°.	172°.
Solubility in	ı light	pe-		
troleum			Appreciably soluble.	Insoluble.
Solubility in	alcohol	•	Practically insoluble in cold; very sparingly soluble in hot.	Sparingly soluble in cold; fairly readily soluble in hot.
Solubility in			Very sparingly soluble.	Readily soluble.
Solubility in sulphide			Very readily soluble.	Sparingly soluble.

FUCOXANTHIN C40H54Og.

This substance was first isolated from fresh brown algae by Willstätter and Page.† It is more difficult to extract this substance from dried algae. Fucoxanthin is a brownish-red substance, which crystallizes from methyl alcohol or light petroleum, and melts at 159:5 to 160:5°. It absorbs iodine

^{*} Ewart: "Proc. Roy. Soc.," 1915 [B], 89, 1. + Willstätter and Page: "Annalen," 1914, 404, 237.

to form a compound $C_{40}H_{54}O_6I_4$. Unlike carotin and xanthophyll, which are neutral substances, fucoxanthin has basic properties, and forms blue salts with hydrochloric and sulphuric acids.

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ANTHOXANTHINS.

FLAVONES AND XANTHONES.

Under the headings of Flavones and Xanthones (two words derived from the Latin and Greek for yellow) are included a number of yellow pigments occurring in the vegetative organs and in the petals of many plants. Owing to their close relationship to the blue colouring matters known as Anthocyanins, Willstätter and Everest,* have proposed the adoption for them of the generic te^{-m}, Anthoxanthin, at first suggested by Marquart in 1835. These yellow pigments are often of considerable economic value as dye-stuffs. They occur naturally in combination with rhamnose or glucose as glucosides and in some cases uncombined, and frequently are also associated with tannins.

The mother substances from which all these substances are derived and from which they derive their name are the two compounds Flavone and Xanthone.

^{&#}x27;Willstätter and Everest: "Annalen," 1913, 401, 189

YELLOW COLOURING MATTERS DERIVED FROM FLAVONE.

There are quite a considerable number of yellow substances occurring in plants derived from flavone, but only a few representative ones will be mentioned here in order to give some idea of the constitution of these compounds.

Chrysin, or dihydroxy-flavone, is a yellow colouring matter occurring in several varieties of poplar, such as Populus nigra and P. pyramidalis.

Quercetin, or tetrahydroxy-flavonol*

occurs with rhamnose in the form of a glucoside in the bark of *Quercus tinctorius*, in the leaves of the horse-chestnut and hop, and in many other plants. Quercetin, in the uncombined

* Flavonol is the hydroxyl derivative of flavone; the relationship between the two substances is shown by the following formulæ:—

state, also is found in the bark of *Pyrus Malus* and in the leaves of *Thea sinensis, Arctostaphylos Uva-ursi, Acacia catechu*, and many other plants.

Rhamnetin, the monomethyl ether of tetrahydroxy-flavonol or quercetin monomethyl ether, occurs in the dried berries of Rhamnus cathartica and R. tinctoria, both of which are used for dveing cotton.

Morin.—This substance, which is isomeric with quercetin, occurs in the wood of Morus tinctoria (yellow wood), where it is accompanied by another colouring matter, maclurin, sometimes called moringatannic acid (see p. 193).

Luteolin.—This is the yellow colouring matter of Reseda luteola, known as "weld"; it is also contained in Genista tinctoria or Dyer's broom.

Other members of this group of substances are Apigenin, occurring in Apium petroselinum, and Fisetin, occurring in Quebracho colorada, and Rhus cotinus or Dyer's sumach.

YELLOW COLOURING MATTERS DERIVED FROM XANTHONE.

There are as yet only three colouring matters known to belong to this group, one of which, euxanthone, does not occur in plants, but in Indian yellow obtained from camel's urine; it has the formula

Gentisin is a yellow colouring matter occurring in Gentiana lutea.

Datiscetin occurring in the form of a glucoside, Datiscin, in Datisea cannabina.

Properties of Anthoxanthins.

- 1. These colouring matters are mostly yellow crystalline solids.
- 2. In water the crystals are hardly soluble, in acids they dissolve readily giving yellow to red solutions, and in alkalies they also are soluble, yielding the same coloured solutions.
- 3. From their solutions they may be precipitated by lead acetate, the precipitate being yellow, orange, or red.
- 4. Aniline or toluidine nitrate and potassium nitrite give a cinnabar red precipitate.
- 5. With ferric chloride a dull green or sometimes a redbrown coloration results.
- 6. On fusion with alkali, decomposition ensues, phloroglucinol and protocatechuic acid being commonly formed, and sometimes resorcinol, resorcylic, or hydroxybenzoic acids,

The solubility of the anthoxanthins in acids is due to the peculiar basic properties of the oxygen atom taking part in the ring formation. The basic nature of the oxygen atom in such circumstances was first observed in the case of the simpler substance pyrone

which dissolves in hydrochloric acid, forming an additive compound of the formula

the oxygen becoming tetravalent. Such additive compounds of anthoxanthins with acids are easily dissociated and do not occur in plants, though it will be seen on page 250 that in the case of the anthocyanins analogous compounds do actually occur naturally.

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ANTHOCYANIN, PHYCOERYTHRIN, AND PHYCOPHAEIN.

Occurring in the cell sap, often in sufficient quantity to mask entirely the green colour of the chlorophyll, are a number of pigments, other than chlorophylls, belonging to various classes of chemical compounds.

Under the collective heading of Anthocyanin are included a number of such pigments of a blue, red, or violet tint occurring in the flowers, fruits, or leaves of many plants.* The first representative of the class to be isolated in a state of purity by Willstätter and Everest† was Cyanin, the blue

^{*}An historical account of our knowledge of these pigments is given by Everest in "Science Progress," 1915, 9, 597.

[†] Willstätter and Everest: "Annalen," 1913, 401, 189.

colouring matter of the cornflower *Centaurea eyanus* and of *Rosa gallica*; closely related to this substance are the anthocyanins of the cranberry (idein), the bilberry (myrtillin), of blue grapes (cenin), of *Delphinium consolida* (delphinin), and of *Pelargonium zonale*, var. *meteor* (pelargonin), *Althaea rosea* (althaein), *Malva sylvestris* (malvin).

The anthocyanins are all glucosides, and on hydrolysis yield one or more molecules of a carbohydrate, together with a so-called anthocyanidin, which compound, unlike the parent substance, is soluble in amyl alcohol. On these facts is based what is known as the *anthocyanin reaction*, according to which a solution of the substance in sulphuric acid yields nothing to amyl alcohol, but after hydrolysis the resulting anthocyanidin may be extracted from the solution quantitatively by amyl alcohol.

The products obtained by the hydrolysis of the various anthocyanins so far investigated are given in the following list:—

```
Cyanin yields Cyanidin
Idæin yields Cyanidin
Enin yields Enidin
Myrtillin yields Myrtillidin
Delphinin yields Delphinidin
Pelargonin yields Pelargonidin + 2 mols. glucose.

+ 1 mol. glucose.
+ 1 mol. glucose.
+ 1 mol. glucose.
+ 2 mols. glucose + 2 mols. p.
hydroxybenzoic acid.
```

The anthocyanidins or non-carbohydrate moiety of the anthocyanins are derivatives of benzo-pyrilium which, as may be seen from the appended formula I.,

is closely related to benzo-pyrone II., the mother substance of the flavones. Both these substances contain a so-called basic oxygen atom which by becoming tetravalent can form additive compounds with acids producing oxonium salts. These salts in the case of the flavones are not stable and do not occur in the plant, but the anthocyanidins yield stable oxonium salts of the type illustrated by the following structural formulæ of a few anthocyanidins:—

It will be seen from these formulæ that the oxygen is tetravalent and that the molecules contain phenolic hydroxyl groups capable of forming salts with alkalies. Moreover, by replacing the chlorine by the hydroxyl group on addition of caustic alkalies the possibility of eliminating water from the molecule arises as follows:—

These considerations explain the colour variations produced by the same cyanidin occurring in the same or in different flowers, it having been found, for example, that the same cyanidin was responsible for the colour of the cornflower and of the rose.*

Thus when combined, as in the case of cyanidin chloride, with mineral acid or in the plant with organic acids, the com-

^{*} Willstätter and Nolan: "Annalen," 1915, 408, 1.

pound has a red tint. When treated with alkali, blue metallic salts are formed, while the arrangement shown in the formula II. represents a neutral compound having a violet tint. The neutral violet tinted delphinin has been isolated from Delphinium consolida by Willstätter and Mieg,* and has been shown to turn blue with alkali, and red with acids; the colour would therefore appear to act as an indicator in the plant itself, showing whether the cell sap is neutral acid or alkaline.

CONNECTION BETWEEN ANTHOCYANINS AND ANTHOXANTHINS

A comparison of the formula of cyanidin chloride on page 250 with that of quercetin on page 245 reveals a close relationship between these two substances, and consequently between the flavones or anthoxanthins and the anthocyanins. Theoretically it should be possible to pass from anthoxanthins to anthocyanins by reduction, or conversely from anthocyanins to anthoxanthins by oxidation. In the plant no doubt this is effected readily enough by enzymes, but in the laboratory it is more difficult, and so far the only transformation effected has been the reduction of quercetin to cyanidin.† These facts provide a confirmation of the views put forward by Wheldale and others previous to this definite experimental evidence. Further evidence for the close relationship between the two classes of compound is provided by the fact that cyanidin is isomeric with luteolin and fisetin while delphinidin is isomeric with quercetin and morin.

EXTRACTION OF ANTHOCYANINS.

The method of extracting anthocyanins varies with the material employed.‡ The method recommended in the case of grape skins is as follows: Extract the skins in the cold with glacial acetic acid and precipitate the dark red filtrate with ether; by heating the deposit so obtained with a solution

^{*} Willstätter and Mieg: "Annalen," 1915, 408, 61. † Willstätter and Mallison: "Sitzungsber, K. Akad, Wiss., Berlin," 1914,

[#] Cf. Willstätter and Mieg: "Annalen," 1915, 408, 61; also Willstatter and Bolton: "Annalen," 1915, 408, 42.

of picric acid a crystalline picrate is formed which separates out on cooling.

ANTHOCYANIN.

The occurrence of a red, blue, or purple pigment, either dissolved in cell sap—the exact colour depending on the acid, alkaline, or neutral reaction of the cell sap—or, less frequently, in the form of needle-shaped crystals, as in the case of *Delphinium ajacis*, is a common phenomenon, and is generally ascribed to the presence of the pigment anthocyanin. It is, however, doubtful whether all such colorations are due to anthocyanins; thus Molisch found that the red colour assumed by the leaves of the aloe, on exposure to high insolation, is due to the formation of carotin within the chloroplasts.

The presence of anthocyanin is due to many causes, light, especially when of high intensity, being important. For example, apples and other fruits and also the vegetative organs of certain plants will not assume a red colour if kept in darkness; on the other hand, light does not appear to be of such importance in the case of the roots of the beet.

In other instances the aerial vegetative organs of many varieties of plants, e.g. certain Chenopodiaceaæ, are characterized by a red colour the presence of which is seemingly independent, or nearly so, of external conditions. Thus Salicornia ramosissima may be found in two forms, one apple green and the other crimson, the intensity of which varies in different years. In such cases there is good reason for supposing that these colours are of an hereditary nature and come true from seed. The same also appears to be true for different forms of beet which are used for horticultural purposes. On the other hand, in the familiar example of the copper beech this is not so, the copper-coloured foliage, due to the combined effect of a red cell sap and the green of the chlorophyll, first originated, it is stated, as a sport and is propagated by means of cuttings.

In the case of anthocyanin in flowers, much more definite information is available owing to numerous investigations upon the inheritance of colour in plants. And although questions relating to genetics are outside our present scope, brief mention

may be made of certain facts in order to illustrate the interrelationship between this branch of botany and chemistry.

In many plants the colour of the flowers depends on various factors:—

C, a chromogen which is not necessarily coloured, and which is, in all probability, a glucosidal flavone.

E, an oxidative enzyme which acts upon C to produce a red colour.

c, another enzyme which acts upon the red pigment, due to the action of E, and further changes it to another pigment so that a different colour results.

A, an antioxidase which inhibits the action of E.

R, a reductase which neutralizes, as it were, the action of E. If a flower, say, of *Lathyrus*, only possesses C or E, then the colour will be white or pale yellow, according to the colour of the chromogen, if present. If the flower with the factor C be crossed with a flower with the factor E, then the colour of the flowers of the offspring will be red, or a deeper colour if e also be present. If either A or R be present, then there will be no difference in the flowers of the offspring as compared with the parents.

By microchemical methods it is possible to ascertain the nature of many of these factors present in the plant, and by such means Keeble and Armstrong* have established many facts relating to the distribution of oxidases in *Primula sinensis* and their relation to the formation of pigments.

The method employed by these authors is the treatment of the tissue with suitable reagents dissolved in media, termed hormones, which solvents cause the plasmatic membranes to become permeable to the reagents, and also renders active the enzymes contained within the cell. One such substance is alcohol.

To take an example: a-naphthol is commonly used as an indicator for oxidases; treatment of the petals of Primula sinensis with an alcoholic solution of a-naphthol decolorizes the tissues, but on then treating the tissue with hydrogen peroxide a coloration obtains, and a comparison of this preparation with the original indicates that the distribution of

^{*} Keeble and Armstrong: "Proc. Roy. Soc., Lond.," B., 1912, 85, 214, 460.

the pigment in the petal coincides exactly with that of a peroxidase. In some varieties of this same plant there is no need to add hydrogen peroxide, the characteristic reaction being obtained by the oxidase reagent alone.

Properties.

The chief physical property of anthocyanin is its absorption spectrum. Engelmann found that it is complementary to that of chlorophyll, the main absorption bands being in the yellow and yellow-green, with minor ones in the blue end of the spectrum.

Questions relating to the energy relationship between this and other pigments and chlorophyll are outside the scope of the present consideration; it may be mentioned, however, that it has been stated that leaves containing anthocyanin have relatively less chlorophyll than those which have no red pigment.

According to Pick and others, anthocyanin is commonly associated with tannins, for a red sap is characteristic of tannin-containing plants, and the precipitate appearing in the palisade cells of *Hydrocharis* on treatment with caffeine and antipyrine closely resembles the precipitates given by the same reagents with tannin. Plants in which this particular pigment does not occur are free from tannin.

The appearance of anthocyanin is closely related to the sugar-content of the tissues in which it occurs.

Ewart* has pointed out that in the case of *Elodea cana-densis* and other aquatic plants the red dye will appear provided the plants be immersed in a weak solution of sugar and exposed to strong sunlight at ordinary temperatures, whilst the red colour does not appear if the plants be grown in water or in diffuse daylight.

These experiments of Ewart were much extended by Overton,† who used *Hydrocharis* and other plants. He found that, in addition to the presence of sugar, light and temperature

^{*} Ewart: "Journ. Linn. Soc., Lond., Bot.," 1895-7, 31, 445; "Ann. Bot.,"

⁺Overton: "Nature," 1899, 59, 296; "Jahrb. Wiss. Bot.," 1899, 33.

were important factors. If the temperature be low, but above freezing-point, then the formation of the red pigment will be promoted, which accounts for the red colour prevalent in alpine plants, since under their conditions of existence sugar tends to accumulate rather than starch. This also is true for arctic plants in which, according to the observations of Wulff,* the leaves are very frequently sugar leaves, and are commonly characterized by the presence of anthocyanin.

In the case of *Hydrocharis* grown in water culture, Overton found that when the temperature and the intensity of light were so balanced that no colour was formed, the addition of 2 per cent of invert sugar caused its appearance in three days. not only in the young leaves but also in the old ones,

Other aquatic plants behaved similarly, but in the case of cut shoots of lilies the red pigment only developed provided sugar were added to the culture solution.

Further experiments showed that the red colour is not formed in those plants, in which the pigment was restricted to the epidermis, when cultivated in sugar solution. Success only resulted in those cases where the colouring matter occurred in the mesophyll.

In view of these facts Overton considered that anthocyanin had some connexion with tannins, and was probably a glucoside (p. 249). A similar view was held by Combes,† who called attention to the facts that, as compared with the green leaves, the red autumnal leaves of Ampelopsis hederacea, etc., contain more sugars and glucosides, the amount of anthocyanin varying directly as the sugars and glucosides; that the dextrins diminish as the sugars and glucosides increase; and that the formation of anthocyanin is not apparently dependent on the insoluble carbohydrates. For these and other reasons he concluded that the substance in question was probably a cyclic glucoside which arose, not at the expense of pre-existent sugars and glucosides nor of chromogens, but in the ordinary course of constructive metabolism; also, he concluded, it was only formed provided that oxygen be present.

The observations of Boodle t also indicate the relationship

^{*} Wulff: "Botanische Beobachtungen aus Spitzbergen," Lund., 1902.

[†] Combes: "Ann. Sci. Nat. Bot.," 1909, 9, 274. ‡ Boodle: "New Phytologist," 1903, 2, 207.

between anthocyanin and sugar. He found that in the leaves of *Rheum*, some of the veins of which had been accidentally severed, anthocyanin made its appearance in the mesophyll supplied by these veins. Boodle then experimented with species of *Oenothera*; all the species examined were not equally responsive, but in the case of *O. biennis* the severance of the midrib at about its middle caused the whole region distal to the cut to become red provided the plant were exposed to daylight. The operation interrupted the path of transport of carbohydrate from the leaf, so that sugar accumulated above the cut, and it is this concentration of soluble carbohydrates which leads to the development of anthocyanin. In this connexion the work of Linsbauer * may be referred to.

That the presence of anthocyanin is connected with nutritive processes there can be no doubt, but other substances besides sugar may come into play; thus Dendy observed that the addition of protein to the water, caused green plants of *Hæmatococcus* to turn red.

Finally, the work of Wheldale[†] on colour inheritance in flowers, points to the conclusion that anthocyanin is a product of the action of an oxidase upon glucosidal flavones, a view which is entirely borne out by the chemical evidence outlined on page 248.

Reactions.

- 1. Soluble in water, alcohol, and ether.
- The solution is coloured according to the reaction, red in the presence of acid and blue when the medium is made alkaline.
 - 3. Strong alkalies decolorize the solution.
 - 4. Basic lead acetate gives a green precipitate.
 - 5. With salts of iron, a green or blue coloration results.

Physiological Significance.

In considering the physiological significance of anthocyanin it must be borne in mind that the substance may occur in almost any organ of a plant, from the root to the flower, and

^{*}Linsbauer: "Oestr. Bot. Zeit.," 1901, 51, 1. +Wheldale: "Proc. Camb. Phil. Soc.," 1909, 15, 137; "Journ. Genet.,"

in plants very remote phyletically one from the other; and that chemically this pigment may not always be exactly the same. Further, as its appearance seemingly depends upon the immediate metabolic condition of the plant, and so in some cases may be sporadic, whilst in other instances it is characteristic of the species or variety or form, care must be exercised in ascribing to it a definite function. Its presence may be due to nothing more than the particular metabolic sequence; in other words, an accident, which, in some examples may be a lucky one for the plant.

It is, of course, not surprising to find that several opinions have been put forward to explain its presence.

According to Pick the dye is a filter to separate from the light entering the leaf certain rays which would be deleterious to the translocation of the starch. Keeble found that in leaves which had the dye on one side but not on the other, the difference in temperature due to the anthocyanin was about 2° C., and he concluded that it may be of value as a protective mechanism against the heating effect of strong sunlight.

Stahl* thought that it absorbs heat and so increases transpiration, especially in the case of tropical plants. Ewart points out that, although this might sometimes be of value, if it were the primary function it would naturally be expected that anthocyanin would absorb the heat rays more particularly. Also Ewart cites his observations on *Elodea* against Stahl's view, and remarks that "since the plants [*Elodea*] are submerged, it cannot possibly be for the purpose of increasing what is non-existent, i.e. transpiration, nor can it perceptibly raise the temperature of a submerged plant". The first argument may no longer be valid, for it appears that a transpiration current may exist in submerged aquatic plants.†

Ewart believes that anthocyanin is to protect the chlorophyll against the action of too strong light. He gives experimental data in support of his view, and cites the observations of Schröder and Klebs to the effect that the pigment is of importance in protecting the chlorophyll in *Hæmatococcus* and the resting spores of many Algæ.

^{*} Stahl: "Ann. Jard. Bot. Buitenzorg," 1896, 13, 137. + See Thoday and Sykes: "Ann. Bot.," 1909, 23, 635-

Ewart does not think that the pigment is an accidental occurrence in all cases, for in *Elodea* it is not formed in diffuse light; on the other hand, in the beetroot it probably has no special function, and may be a waste product of metabolism.

Stahl also expressed the view that the red colour may protect the plant from the predatory tendencies of herbivorous animals. Such a function is, of course, quite secondary, but may be of use to the plant when the animals have an antipathy to certain hues. Tichler considered that anthocyanin promotes the anabolic processes of the plant.

Overton agrees with the view of Stahl that the presence of anthocyanin may promote nutritive processes by the absorption of heat.

According to Buscalioni and Polacci* anthocyanins may increase the osmotic forces of the cell, but they are careful to point out that they may perform many functions in different plants.

Wulff considers that the pigment is of value in the absorption of extra radiant energy, and is of great importance in arctic plants, for instance, which live under conditions unfavourable for metabolic activities.

Combes holds views similar to those of Palladin, that anthocyanin is closely connected with respiration. If the sugar content increases the rate of respiration is accelerated, and this leads to the formation of the pigment.

It may be remarked that most of the above opinions were put forward before the work of Palladin on respiration and the relationships between pigments and enzymic activity appeared. And, in view of this, some of the earlier experiments appear to require reconsideration from Palladin's point of view.

PHYCOERYTHRIN.

Phycoerythrin is a red pigment commonly occurring in red sea-weeds, especially when growing in deep water. It has recently been investigated by Hanson,† on whose account the following description is based:—

Phycoerythrin is easily soluble in water, giving a rose-

^{*} Buscalioni and Polacci: "Atti. Inst. Bot., Pavia," 1904, II, 8, 1, 135, † Hanson: "New Phytologist," 1909, 8, 337.

coloured solution which exhibits a well-marked orange fluoresence; the spectrum shows that the chief absorption is that of the blue-green rays.

Preparation.

To prepare a solution of phycoerythrin the red sea-weed, *Ceramium rubrum*, which is one of the best to use, is washed in ordinary water to free it from sea salts and adhering sand. It is then soaked in distilled water; in two days most of the pigment will have diffused out. The solution is filtered through glass wool and a few drops of eucalyptus oil added as an antiseptic, for putrefaction soon sets in.

It is a matter of great difficulty to obtain a pure sample of phycoerythrin, for, in an aqueous solution, it passes over into an irreversible gel,* even when kept at o° C. This, of course, renders ordinary filtration extraordinarily slow, and thus increases the difficulty of purification.

The solid phycoerythrin may be prepared from the aqueous solution by concentrating it under reduced pressure at a temperature not higher than 38° C.; any precipitate which comes down during this process must be filtered off. Methylated spirit is then added to the concentrated solution until the fluorescence disappears. The precipitated phycoerythrin is allowed to settle and the more or less clear supernatant fluid is filtered off, again treated with alcohol, and filtered. The operation is repeated until the red colour has entirely disappeared from the solution. The precipitates are washed by decantation with 70 per cent alcohol; the pigment, in a pasty mass, is placed in a clock glass and dried in a vacuum.

Reactions.

The following reactions are among those recorded by Hanson:—

- I. Phycoerythrin is precipitated from its solution by alcohol, by small quantities of mercuric chloride, and by saturation with ammonium sulphate and magnesium sulphate.
- 2. When dilute acids are added gradually, the fluorescence first disappears, leaving a somewhat opalescent solution of

a lilac-pink tint. After the lapse of two days a pink precipitate comes down.

- 3. Ammonium hydrate in small quantities removes the fluorescence; in excess, a yellowish-brown coloration results.
- 4. Caustic soda or potash in small quantities causes the red colour to disappear, the solution turning opalescent and yellowish-brown in colour; on standing, a brownish precipitate comes down.
- 5. The solution is immediately decolorized by bleaching powder, bromine water or a solution of iodine in potassium iodide.
- 6. Mercuric chloride solution in small quantities gives a lilac-grey precipitate, the solution then being yellowish in colour.
 - 7. Ferric chloride gives a pinkish-brown precipitate.
- 8. Boiled with nitric acid a yellow colour results which turns to orange on adding an excess of ammonia.
 - 9. Boiled with Millon's reagent a deep red colour results.
- 10. The addition of a caustic soda solution followed by a drop or two of dilute copper sulphate gives a greenish tint.
- 11. Digestion with pepsin, in the presence of hydrochloric acid, has no result.
- 12. On digestion with trypsin in the presence of sodium carbonate, the phycoerythrin loses its colour, and the solution contains a very small amount of leucin, but no tyrosin.
- 13. On hydrolysis with acids, tyrosin is found in very small amounts, but leucin occurs in greater quantities.

From these and other facts it is concluded that phycoerythrin is a colloidal nitrogenous substance allied to the proteins; it is not a true protein, since its nitrogen content is too low and it does not give the biuret reaction. It is impossible to say anything more definite regarding its chemical nature until it has been prepared in a pure state in quantities sufficient for analysis.

Physiologically, phycoerythrin acts as a pigment complementary to chlorophyll. It absorbs the blue-green rays, and degrades them to yellow and red light of just those wavelengths which the chlorophyll can absorb.

PHYCOPHAEIN.

As is well known, a brown colouring matter may be extracted by water from the Phæophyceæ and other brown

Algæ. Hitherto this has generally been considered to be due to the presence within the cells of a definite colouring matter of a protein nature. According, however, to the work of Molisch * and Tswett,† this is not the case. The brown colouring matter is really due to post-mortem changes, the oxidation of a water soluble chromogen. An extract prepared with distilled water is at first colourless, but will turn yellow if the solution is made alkaline in reaction, e.g. by tap water, and finally brown owing to oxidation. If the reaction be made acid decolorization will result. With regard to the chemistry of this substance little, if anything, is known.

Tswett finds that the natural colour of brown sea-weeds is due to the presence of several pigments in the chloroplast, amongst which is carotin, but until more information regarding the relationships between these and similar pigments is available, it is hardly profitable to consider them further.

With regard to the physiological significance of these pigments in the Algæ, the work of Gaidukov‡ on complementary chromatic adaptation may be consulted.

Although it is not proposed to enter into a detailed consideration of the phenomena of respiration here, brief mention may be made of Palladin's § conceptions on the subject on account of the rôle he ascribes to colouring matters and allied substances in respiratory activity.

Occurring in plants are pro-chromogens which may be glucosides or may be decomposition products of proteins. These pro-chromogens, by the action of enzymes, give origin to chromogens.

Chromogens are widely distributed in the vegetable kingdom, in fact are universally present in those parts of plants which are respiring; they, however, vary in amount at different seasons of the year and according to the physiological condition of the plant. For instance, in the spring they occur in abundance in the young leaves, and in the autumn the old and dead leaves also contain much owing to the lack of coordination of enzymic activity.

^{*} Molisch: "Bot. Ztg.," 1894, 52, 181; 1895, 53, 131; 1905, 63, 131.

[†] Tswett: "Ber. deut. bot. Gesells.," 1906, 24, 235. ; See Blackman: "New Phytologist," 1904, 3, 237.

[§] Palladin: "Ber. deut. bot. Gesells.," 1908, 26a, 125, 378, 389; 1909, 27, 110.

At other times the amount of chromogens is not very great, but may be increased by suitable treatment. Thus Palladin found that leaves kept for a week in a strong solution, 20 to 30 per cent, of cane sugar showed a great increase, whereas leaves kept in distilled water and also untreated leaves of the plant showed no such increase. A bright illumination also increases the amount of chromogens.

The chromogens are acted upon by oxidases in the presence of oxygen and yield pigments which may be reduced by reducing enzymes or reductases. Carotin and Xanthophyll provide convenient examples. Carotin, $C_{40}H_{56}$, is acted upon by an oxidase and converted into Xanthophyll, $C_{40}H_{56}O_2$, which in turn is acted upon by a reductase yielding carotin. This action is comparable to that of the hæmoglobin in the blood, and in fact Palladin has termed all such respiratory pigments of plants, no matter what their composition may be, phytohæmatins.

It does not follow that all definite coloured compounds are formed during respiration; it all depends on the relative activities of the oxidases and the reductases. A pigment will make its appearance provided the oxidases are the more potent, but if the reductases are the more active no pigment will appear.

The method of indicating the presence of a chromogen is obvious; the material to be examined is extracted and heated to a degree of temperature sufficient to destroy any enzymes present. To this preparation is added peroxidase and hydrogen peroxide; if a chromogen were present originally, then a coloration will result, usually brown, red, or purple.

SECTION VI.

NITROGEN BASES.

Ammonia is said to have basic properties because it can form salts by combining with acids. This salt formation, which may be illustrated by the conversion of ammonia into ammonium chloride, is due to the unsaturated nature of the trivalent nitrogen atom, and its tendency to assume the pentavalent condition.

$$N^{iii} \begin{array}{c} H \\ H \end{array} \rightarrow \begin{array}{c} N^v \end{array} \begin{array}{c} H \\ H \\ H \\ H \end{array}$$

Ammonia Ammonium chloride

The replacement of one or more of the hydrogen atoms in ammonia by organic radicles, such as methyl, CH_3 —, ethyl, C_2H_5 —, or phenyl, C_6H_5 —, gives rise to compounds known as amines or substituted ammonias, which still retain the property of salt formation possessed by the parent substance ammonia.

For example:-

$$CH_{3}N^{iji} \bigg\langle \begin{matrix} H \\ + \ HCI \end{matrix} \ \Rightarrow \ CH_{3} \!\!-\!\! N^{\nu} \ \bigg\langle \begin{matrix} H \\ H \\ H \end{matrix} \bigg\rangle$$

Methylamine

Methylamine hydrochloride, or Methyl ammonium chloride

$$(C_2H_5)_2 = NH + HI \rightarrow (C_2H_5)_2 = N - H$$

Diethylamine

Diethylamine hydriodide

$$(CH_3)_3 \equiv N + HBr \rightarrow (CH_3)_3 \equiv N$$

Representation of the second secon

Trimethylamine

Trimethylamine hydrobromide

These three substances, CH_3NH_2 , methylamine, $(C_2H_5)_2NH$, diethylamine, and $(C_2H_5)_3$; N, triethylamine, are types of three different classes of amines, known respectively as primary,

secondary, and tertiary amines, according as one, two, or three of the hydrogens of ammonia have been replaced by organic radicles.

Tertiary amines are also known in which the nitrogen atom takes part in the formation of a ring, as, for example, in pyridine—

which may be regarded as being derived from ammonia by the replacement of three atoms of hydrogen by the five carbon ring—

Pyridine, being a substituted ammonia, can form salts by changing the valency of its nitrogen atom from three to five, as follows:—

Pyridine Pyridine hydrochloride

Secondary amines containing a nitrogen atom in the ring are also known.

Thus, when pyridine is reduced by nascent hydrogen, six atoms of hydrogen are added on, and a substance known as piperidine is produced; this substance is a secondary amine, since it now has a hydrogen atom attached to its nitrogen. Like pyridine, it can also form a salt with hydrochloric acid.

$$\begin{array}{cccc} \operatorname{CH}_2 & \operatorname{CH}_2 \\ \operatorname{CH}_2 & \operatorname{CH}_2 & \operatorname{CH}_2 \\ \operatorname{CH}_2 & \operatorname{CH}_2 & \rightarrow & \operatorname{CH}_2 \\ \operatorname{CH}_2 & \operatorname{CH}_2 & \rightarrow & \operatorname{CH}_2 \\ \operatorname{NH} & & \operatorname{N} - \operatorname{H} \\ & & & \operatorname{H} & \operatorname{CI} \\ \end{array}$$

From the above examples it will be seen that the presence of a trivalent nitrogen atom in a compound, whether in a ring or attached to a straight chain, will, as a rule, confer on that compound basic properties, owing to the tendency of that nitrogen to become pentavalent by combining with an acid and producing a salt. It is this property which gives rise to the term *Nitrogen base*.

The discovery and isolation from natural sources of a number of nitrogen bases, such as cinchonine, quinine, brucine, strychnine, morphine, etc., having properties analogous to those of the alkalis in being able to form salts with acids, led to their designation as alkaloids or alkali-resembling substances. As the number of such substances increased, a distinction began to be made between animal and vegetable alkaloids. The term alkaloid is, however, better reserved for nitrogen bases of vegetable origin; it was at one time suggested that the term should include only derivatives of pyridine, quinoline, and isoquinoline.

but this definition excludes such compounds as stachydrine and hygrine, etc., which are pyrrolidine derivatives, and also the purine bases which, according to most authors, should be included among the alkaloids.

This difficulty is, however, overcome by defining alkaloids as nitrogen bases of vegetable origin whose nitrogen atom forms part of a ring.

Even this definition is not entirely satisfactory, as it would include substances which, owing to their properties, could hardly be classed as alkaloids, and excludes others, such, for example, as hordenine.

ALKALOIDS. Occurrence.

The alkaloids do not appear to have a wide distribution in the vegetable kingdom. Amongst the Angiosperms, the Apocynaceæ, Leguminosæ, Papaveraceæ, Ranunculaceæ, Rubiaceæ and Solanaceæ stand out in the provision of several of these substances. The Labiatæ, Rosaceæ, Orchidaceæ, and Monocotyledons and Gymnosperms very rarely contain them.

Alkaloids may occur in solution in the cell sap, especially in young parenchyma: in older tissues the substances in question may be stored in the solid state. They are found in the seeds and fruits more particularly, but in the case of the alkaloids of the Solanaceæ and some other plants they occur in the leaves, whilst the roots are the chief sources of the alkaloids of Aconitum, Corydalis, and Hydrastis. The cinchona alkaloids, and also pelletierine of the pomegranate, are contained in the bark of their respective trees.

Classification.

The classification of the alkaloids is based upon the structure of the nucleus upon which their molecules are built up. Five groups of alkaloids are accordingly recognized.

I. Pyridine Alkaloids.—These, as the name implies, are all derivatives of pyridine, and include:—

Conjine from Conjum maculatum.

Arecolin from Areca catechu.

Trigonellin from Trigonellum fænum, Pisum sativum, etc.

Piperine from Piper, and

Nicotine from Nicotiana tabacum.

Some idea of the structure of the molecules of alkaloids belonging to this group may be obtained from the two following constitutional formulæ, which represent conline and nicotine respectively:—

From these formulæ it may be seen that coniine is derived from pyridine, or more strictly from piperidine—

whilst nicotine contains two rings, one a pyridine ring and the other a pyrrolidine ring—

such as is also found in proline (see p. 316).

II. Pyrrolidine Alkaloids.—This is a small group, comprising as yet only three alkaloids, namely:—

Hygrine and Kuskhygrine, from the leaves of *Erythroxylon* Coca. and

Stachydrine, from the tubers of *Stachys tuberifera** and leaves of *Citrus vulgaris.*†

The constitution of stachydrine is as follows:-

$$\begin{array}{c|c} CH_2-CH_2\\ & \downarrow \\ CO-CH & CH_2\\ \downarrow \\ O-N(CH_3)_2\\ Stachydrine \end{array}$$

showing it to be a dimethyl betaine of pyrrolidine.

III. Tropane Alkaloids.—The alkaloids belonging to this group are derivatives of tropane—

which substance, as may be seen, contains both a six-membered piperidine ring and a five-membered pyrrolidine ring.

The group includes alkaloids from the four Natural Orders:—

Solanaceæ, e.g. Atropine, Hyoscine, Hyoscyamine.

* Planta and Schulze: "Arch. d. Pharm.," 1893, 305; "Ber. deut. chem. Gesells.," 1893, 26, 939; Schulze and Trier: "Ber. deut. chem. Gesells.," 1909, 42, 4654; "Zeit. physiol. Chem.," 1910, 67, 59.

⁺ Jahns: "Ber. deut. chem. Gesells.," 1896, 29, 2065.

[#] For an explanation of the term betaine, see p. 265.

Erythroxylaceæ, e.g. Coca alkaloids, such as Cocaine and Tropacocaine.

Myrtaceæ: Pelletierine, Isopelletierine, etc., from *Punica* granatum (pomegranate).

Papilionaceæ: Cytisine from Cytisus Laburnum; Lupinine from Lupinus luteus and Lupinus niger.

Most of the above alkaloids have a very complex constitution, and the formula of only one will be given, namely, cocaine:—

- IV. Quinoline Alkaloids.—These fall into two groups:—
- (a) Cinchona alkaloids, such as Quinine, Cinchonine, etc., from the bark of various species of Cinchona (Rubiaceæ).
- (b) Strychnos alkaloids, such as Strychnine and Brucine from Strychnos nux vomica, S. Ignatii, etc., and Curarine from Strychnos toxifera (Loganiaceæ).

The constitution of quinine is represented by the following formula *:—

$$CH_{2} - CH$$

$$CH_{2} - CH - CH = CH_{2}$$

$$CH_{2} - CH_{2}$$

$$CHOH - CH - N$$

Quinine

from which it will be seen to contain a quinoline ring.

 * This formula, though probably correct, has not yet been confirmed by synthesis.

The constitution of strychnine and brucine has not yet been determined, though possible formulæ have been suggested by Perkin and Robinson.*

- V. Isoquinoline Alkaloids.—These may be divided into the three following groups:—
 - (a) Papaverine group, including Papaverine, Narcotine, Laudanosine, etc., closely allied to which are Hydrastine and Hydrastinine from Hydrastis canadensis.
 - (b) Morphine group, including Morphine, Apomorphine, Thebaine, and Codeine.
 - (c) Berberine group, including Berberine and Corydalis Alkaloids.

The constitutional formulæ for alkaloids of this group are for the most part exceedingly complex, and it will suffice here merely to show the skeleton formulæ of a member of each group:—

Papaverine Morphine + Berberine

In addition to the alkaloids mentioned above, there are a very large number which cannot as yet be classified, since their constitution is still unknown; these include amongst others ergotinine from ergot, colchicine from Colchicum, taxine from Taxus baccata, aconitine from Aconitum Napellus, delphinine from Delphinium, etc.

GENERAL PROPERTIES OF ALKALOIDS.

The alkaloids are, as a rule, composed of the four elements, carbon, hydrogen, nitrogen, and oxygen, but a few are known, such as coniine, nicotine, and one or two little-known ones, such as hymenodictine and conessine (from bark of Wrightia antidysenterica), which contain no oxygen.

^{*} Perkin and Robinson: "J. Chem. Soc., Lond.," 1910, 97, 305. † This formula is subject to revision.

There are a few alkaloids which are liquid, e.g., coniine, nicotine, pelletierine, sparteine, etc., but by far the greater number are colourless crystalline solids. They are, as a rule, insoluble in water, but dissolve in neutral organic solvents. such as ether, amyl alcohol, chloroform, carbon tetrachloride, etc., whereas their salts have just the opposite solubilities.

They are mostly free from smell, but conline, nicotine, and sparteine have strong odours.

Most of them have a bitter taste and are possessed of marked physiological or toxic properties.

They are all bases, and accordingly have an alkaline reaction in solution, though it must be borne in mind that aqueous solutions of the salts usually have a strongly acid reaction due to hydrolytic dissociation.

The majority of alkaloids are optically active, rotating the plane of polarized light to the left, though a few, such as coniine, laudanosine, pelletierine and pilocarpine, are dextrorotatory.

GENERAL REACTIONS OF ALKALOIDS.

The alkaloids are precipitated from solution by a large number of different reagents with formation of amorphous or sometimes crystalline precipitates.

The commonest of these reagents are the following:-

- I. A solution of iodine in potassium iodide, sometimes known potassium ter-iodide, gives a chocolate-brown precipitate.
- 2. Mercuric iodide in potassium all of which iodide.
- 3. Tannic acid,
- 4. Phosphotungstic acid.
- 5. Auric chloride,
- 6. Platinic chloride (see p. 276),

give colourless amorphous precipi-

which give crystalline precipitates often having characteristic

The alkaloids are, however, not the only substances which are thrown out of solution by these reagents, since most nitrogen bases behave in a similar way, and the formation of a precipitate is therefore not conclusive proof of the presence

of alkaloids. On the other hand, if none of the above reagents produce precipitates, it is tolerably certain that there are no alkaloids present.

In examining plant tissues for alkaloids, Errera recommends testing the fresh sections with alkaloidal reagents and also sections which have been soaked in a five per cent alcoholic solution of tartaric acid. In the second case no precipitate should be obtained, owing to the extraction of the alkaloid.

The final identification of the various alkaloids is usually effected by means of colour reactions.

Thus, if a section of the endosperm of *Strychnos nux vomica* be mounted in a few drops of strong sulphuric acid, the presence of strychnine is indicated by a red coloration of the cell-contents. This colour will change to violet on placing a small crystal of potassium chromate beneath the cover-glass.

Similarly, a section of the rhizome of Aconitum Napellus, when treated with a few drops of 50 per cent sulphuric acid, will show a carmine red coloration, due to the presence of aconitine, in the parenchyma surrounding the vascular bundles. This reaction is the more marked when the section has been previously warmed in a solution of sucrose.

These colour reactions are very numerous; for them the larger text-books and monographs must be consulted.

Isolation.

Most alkaloids do not occur free in the plant, but combined with some acid in the form of a salt; the acids most commonly met with are tannic, malic, citric, succinic and oxalic, while acetic and lactic acids are rarer; some acids occur only in connexion with certain alkaloids, such as meconic acid with opium and quinic acid with quinine.

In some few cases the alkaloids can be extracted from their natural sources by means of organic solvents, such as chloroform, carbon tetrachloride, ether, etc., but in the majority of cases the alkaloid requires to be set free first by the addition of an alkali, such as lime or baryta, since only the free bases, and not the salts, are soluble in the abovementioned solvents.

The material to be extracted is mixed with slaked lime

and carefully dried, and then extracted in a Soxhlet extractor with chloroform or carbon tetrachloride; the extract is then shaken up with dilute sulphuric acid, whereby the sulphate is formed; the acid layer containing the salt in solution is then run off and evaporated, when the alkaloid salt crystallizes out and can be further purified by recrystallization.

Example.—Preparation of quinine from cinchona bark. Twenty grams of quicklime are stirred up with 200 c.c. of water and then thoroughly mixed in a mortar with 100 grams of cinchona bark which have been ground up in a coffee mill. The resulting mixture is then dried over a water bath, care being taken to prevent the formation of lumps. The dried substance is then extracted in a Soxhlet apparatus with chloroform. The extract is then shaken up with 25 c.c. of dilute sulphuric acid, the chloroform layer being run off from below; it is then shaken up with water several times and the water and acid extracts are mixed together and neutralized with ammonia. On evaporating the solution, quinine sulphate crystallizes out; the amount obtained rarely exceeds 1-2 grams in weight.

N.B.—A rapid way of testing a piece of bark for quinine consists in heating it in a dry test tube. If there is any quinine present, the bark will give off a carmine-coloured vapour.

THE ORIGIN OF ALKALOIDS IN THE PLANT.

According to Pictet,* alkaloids are produced in the plant in two successive stages, involving (I) the breakdown of complex nitrogenous substances, such as protein or chlorophyll, with the production of relatively simple basic substances; (2) the condensation of these relatively simple substances with other compounds present in the plant, with the formation of the complex molecules possessed by the alkaloids.

The processes of metabolism within the plant would therefore be strictly analogous to those taking place in the animal body, in which waste products, such as phenol, glycine, etc., are coupled up with other substances, such as sulphuric or benzoic acid, before being eliminated.

^{*} Pictet: "Arch. Sci. Phys. Nat.," 1905, [iv], 19, 329; "Ber. deut. chem. Gesells.," 1907, 40, 771.

Pictet is further of opinion that among the commonest changes within the plant are the methylation of hydroxyl or amino groups by formaldehyde, according to the equations—

$$ROH + CH2O = ROCH3 + O$$
and
$$RNH + CH2O = RNCH3 + O$$

the resulting methylated compounds being then able to undergo intramolecular transformation, by which the methyl group can enter the ring, and so produce, for example, a pyridine ring from methyl pyrrole, a reaction which he has been able to effect in the laboratory by heat.

Similar changes would also explain the formation of quinoline and isoquinoline, and it thus becomes possible to account for the origin of the pyridine and quinoline rings which occur in alkaloids, by assuming them to have been produced as above from pyrrole or indole rings, which are the normal constituents of protein (e.g. proline, histidine, tryptophane, etc.).

In support of these views, Pictet states that he was able to isolate by steam distillation from various leaves,* etc., treated with sodium carbonate, a number of simple bases which he calls proto-alkaloids; these include pyrrolidine and methyl pyrroline—

whose origin from the protein molecule is readily intelligible, in view of the fact that a similar ring occurs in proline, the cleavage product of a number of proteins. It is assumed that these proto-alkaloids are subsequently methylated, rearranged and condensed as described above to form the more complex alkaloids.

It has been suggested by Pictet that the secretion of

^{*} The leaves used were those of tobacco, carrot, parsley, and coco.

alkaloids by plants is merely due to the inability of such plants to get rid of their nitrogenous products of metabolism by any other means than by converting them into alkaloids, which, though poisonous to animals, are not toxic to the plants themselves.

PTOMAINES.

Associated with the simplest form of plant life, namely, bacteria, a number of different basic substances are found, some of very simple constitution, such as methylamine, CH_3NH_2 , dimethylamine, $(CH_3)_2NH$, trimethylamine, $(CH_3)_3N$, putrescine, $NH_2(CH_2)_4NH_2$, cadaverine, $NH_2(CH_2)_5NH_2$, and others rather more complex, such as choline, muscarine, neurine, collidine, etc., and some of unknown constitution, such as mydaleine and sepsine. These substances are known as ptomaines,* from the fact that they are usually associated with decomposing flesh; some of them, such as putrescine and cadaverine, are practically non-poisonous, while others are highly toxic, producing increased salivation, diarrhæa, vomiting, etc.

On the whole, however, it is at least doubtful whether the manifestations of ptomaine poisoning are to be attributed entirely to these substances; it would seem more likely that they were largely due to bacterial toxins, a class of substance related to the albumoses, which have the power of inducing the formation in the blood of antibodies, or, as they are better called, anti-toxins. Similar toxins or toxalbumins also occur in certain of the higher plants, as, for example, abrin, obtained from *Abrus precatorius*, and ricin, which occurs in *Ricinus*.

The so-called ptomaines are all decomposition products of the complex nitrogenous substrate upon which the moulds or bacteria are growing, but are not actually found within the organisms themselves.

In the higher forms of plant life, on the other hand, these bases are actually secreted by and stored up in the plants; the substance muscarine, for example, occurring in the fungus † Amanita muscaria.

^{*} From the Greek word πτωμα, meaning corpse.

[†] The same remark, of course, applies also to the Angiosperms, which contain their alkaloids stored up in different parts of their structure.

Considerations of space will not permit more than the very briefest reference to the chemistry of these substances.

The compounds choline, muscarine, betaine, and neurine are closely related, as may be seen from their formulæ:—

the relationship to each other of the first three being that of alcohol, aldehyde and acid anhydride.*

Choline and muscarine occur in the toad-stool, Amanita muscaria. Betaine and choline frequently occur together, as for example in the germ of Hordeum sativum, Triticum sativum, Vicia sativa, Lathyrus sativus, Gossypium herbaceum, and several other plants. Betaine alone occurs in the juice of the beet † and in tubers of Helianthus tuberosus. Choline is far more widely distributed, and occurs in seeds and fruits of a very large number of plants, such as Pinus cembra, Areca Catechu (nut), Cocos nucifera (endosperm), Acorus calamus (root), Fagus silvatica, Cannabis sativa and C. indica, Humulus Lupulus, etc.

Neurine does not occur in plants, but is produced in putrefying fish and meat. Muscarine and neurine are both very poisonous, whereas choline is comparatively innocuous.

All these substances are strong bases, and answer the general reactions for alkaloids (which see).

A few other bases of comparatively simple constitution which occur in plants may here be mentioned.

Trimethylamine, (CH₃)₈N, is a very volatile substance which occurs in the seeds of *Mercurialis annua* and in the flowers of *Cratægus Oxyacantha*, *Pyrus Aucuparia*, and many other plants,

* The name betaine is derived from the fact that this substance was first obtained from the beetroot (Beta vulgaris). It is the anhydride of hydroxytrimethylamino-acetic acid.

The alkaloid stachydrine (see p. 267) is a derivative of this substance.

† For the preparation of betaine from this source, see "Ber. deut. chem. Gesells.," 1912, 45, 2411.

and is given off from the leaves of *Chenopolium Vulvaria*. It is also readily produced from choline and betaine, and is, therefore, commonly produced from putrifying animal or vegetable matter containing lecithin (see p. 48).

Parahydroxyphenylethylamine, HO CH₂CH₂NH₂, is a substance occurring in ergot, which has a marked pressor action on the circulation, and causes contraction of the uterus. Its close relationship to tyrosine, from which it can be obtained by loss of carbon dioxide, is of interest.

Hordenine, $HO \searrow CH_2CH_2N(CH_3)_2$, is the dimethyl derivative of the previous compound, and occurs in barley.

The fact that all nitrogenous bases form crystalline derivatives with such substances as platinic or auric chlorides, or with picric or picrolonic acids is frequently made use of for isolating or identifying small quantities of these substances (see choline, lecithine, p. 48); since the derivatives produced can, as a rule, be identified by their crystalline form and melting point, they provide a certain method of recognizing substances which do not give any characteristic colour reactions.

An additional advantage of the method lies in the fact that the reagents employed (auric or platinic chloride, etc.) being substances of high molecular weight produce crystalline derivatives whose weight is very considerably greater than that of the substance which is being isolated, and thus ponderable quantities of substance may be obtained from comparatively small amounts of material.

PURINE BASES.

Under this heading are included such substances as caffeine, theobromine, xanthine, guanine, etc., which are called purine bases because they are all derivatives of the same substance, purine, whose formula is given below:—

This substance, which is also the mother substance of uric acid, does not occur in nature, but has been synthesized by Fischer.

By writing the formula somewhat differently, as follows:-

it will be seen that it is composed of two rings, the upper one, which is six membered, being a so-called pyrimidine ring, while the lower one, which is five membered, is an imidazol or glyoxaline ring, the same as occurs in histidine (see p. 316).

The relationship between purine, xanthine, theobromine and caffeine is best understood from the following considerations.

Xanthine may be regarded as purine with the addition of two atoms of oxygen attached to the carbon atoms numbered 2 and 6; and it is accordingly called 2:6 dioxypurine, and is given the formula:—

Xanthine or 2:6 dioxypurine

From this compound theobromine and caffeine are derived by replacing two and three atoms of hydrogen respectively by methyl groups, as may be seen from the following formulæ:—

Xanthine is widely distributed among plants, notably in sprouting seedlings, and occurs also in tea leaves and in the juice of the beetroot.

Theobromine occurs chiefly in the fruit of *Theobroma Cacao* (1.5-2.4 per cent), and a small quantity also occurs in kola nut and in tea leaves, but not in coffee; it acts as a powerful diuretic and has a stimulating effect on the central nervous system, but is less powerful in this respect than caffeine.

Caffeine occurs to the extent of about 1-2 per cent in kola nuts, '1-'8 per cent in cocoa beans, from 2-5 per cent in tea leaves, from 0'8-1'7 per cent in coffee beans, and from 2'5-3 per cent in the fruit of *Paullinia cupana*; the latter substance ground up into a paste is consumed in South America under the name of guarana. The so-called Maté or Paraguay tea, the dried leaves of *Ilex paraguensis*, contains about 0'2-1'6 per cent of caffeine.

Caffeine is a powerful cerebral stimulant, but also acts somewhat on the heart; it is furthermore a powerful diuretic.

Three further purine bases deserve mention, namely, Adenine, Hypoxanthine and Guanine, the formulæ of which are as follows:—



All three substances have been obtained by the hydrolysis of nucleo-proteins from plants (see p. 326) and of nucleic acids from yeast * and from *Triticum sativum.*†

^{*} Schittenhelm and Schröter: "Zeit, physiol. Chem.," 1904, 41, 290. + Osborne and Harris: id., 1902, 36, 85; Osborne: "Amer. Journ. Pharm.," 1903, 9, 69.

Guanine and Hypoxanthine are usually found together; they occur in sprouting seeds of a number of plants, notably Cucurbita Pepo, Acer pseudoplatanus, Vicia sativa, Trifolium pratense, Lupinus luteus, Hordeum sativum, and in the juice of the beet, etc.

Adenine, which is less widely distributed, likewise occurs in the juice of the beet and in tea leaves, and has also been found in leaves of *Trifolium repens*.

Uric acid, which is systematically named 2:6:8 trioxy-purine, has the formula—

It does not occur in plants, but is a well-known product of metabolism in the animal world. In view of the close relationship between this substance and the other purine bases, the assumption does not seem unwarranted that the purine bases in the plant are also waste products (see below). And, in this connexion, it is interesting to find that the presence of urea, in very small amounts, has been observed by Fosse* in the higher plants. To what extent this substance is a physiological product of the cell is doubtful.

The identification of individual members of the purine bases is not very easy, although the recognition of a purine base as such is rendered simple by the so-called murexide test which is given by practically all the members of this group of compounds.

The test consists in evaporating the substance (uric acid or caffeine may be used) in a porcelain basin with dilute nitric acid over a water bath. A yellowish residue remains which on the addition of ammonia or by exposure to ammonia vapour turns pink; potash changes the colour to purple.

The identification of caffeine in plants has been the subject of numerous researches †; it is precipitated by several alkaloidal reagents from solutions containing concentrated hydrochloric

^{*} Fosse: "Compt. rend.," 1912, 155, 851; 1913, 156, 567. See also page 340. + Clautriau: "Nature et Signification des Alcaloïdes végétaux," Brussels, 1000,

acid, but not from neutral solutions; these precipitates are, however, not characteristic. Behrens* has described methods of identifying this substance with the help of mercuric chloride and of silver nitrate and nitric acid. The method is as follows:—

Fifty mgs. of dried tea leaves are coarsely powdered and mixed with quicklime and sufficient water to make a crumbly mass. The mixture is then dried and extracted with alcohol; the extract is evaporated drop by drop on a microscope slide and finally the residue is sublimed by heating until it turns brown, the vapour being condensed on a second slide held about 2 mm. above it. The sublimate consists of well-formed needle-shaped crystals. A drop of water containing a trace of hydrochloric acid is then placed near the sublimate and a grain of mercuric chloride is dissolved in the drop. On drawing the mercuric chloride solution through the sublimate, colourless glistening prismatic crystals are produced.

Silver nitrate in the presence of a small quantity of nitric acid produces under similar circumstances woolly aggregates.

PHYSIOLOGICAL SIGNIFICANCE OF NITROGEN BASES.

In considering the physiological significance of alkaloids, questions naturally arise with regard to their place in the metabolism of the plant. Are they connected with the elaboration of food? or are they so much waste material, bye-products of metabolism, corresponding to uric acid and such-like substances excreted by the higher animals? Unfortunately, definite answers are not possible; what may be true of one group of nitrogen bases may be incorrect for another, and in any case the answers would not appear to be of general application, owing to the restricted occurrence of some of these compounds in the vegetable kingdom.

Certain organisms, more especially lower ones, can use alkaloids as a raw food-material, provided they be supplied in a sufficiently dilute state; thus certain Fungi seemingly can assimilate morphia. Amongst the Algae, Comère † found that *Ulothrix subtilis* and *Spiregyra crassa*, grown under aseptic

^{*} Behrens: "Anleitungen z. mikrochemischen Analyse d. wichtigsten organ. Verbindungen," 1897, IV, 14.

[†] Comère: "Bull. Soc. Bot., France," 1910, 57, 277.

conditions and in a solution free from nitrates, could make use of certain alkaloids as a source of nitrogen. Of the alkaloids used, this was found to be true for the sulphates and hydrochlorides of atropine, cocaine and morphine; quinine, although it had no deleterious action, was not assimilated, whilst strychnine showed a marked toxic action. Clautriau* found that alkaloids supplied to the higher plant as the sole source of nitrogen are not utilized.

With regard to the higher plants, De Vries considers that alkaloids are not essential for the well-being of the plant, since in the germination of the seed of the potato, the thornapple (Datura Stramonium) and nux vomica (Strychnos nux vomica), little or no diminution in the substances in question occurs. This opinion is to a certain extent supported by the fact that their presence depends, at any rate in some cases, on the conditions of cultivation; for instance, quinine does not occur in cinchona cultivated in hot-houses in this country.

Lotsy † considers that alkaloids, such as quinine, are not decomposition products of proteins, but direct synthetic substances. In the case of *Cinchona*, he found that the bases occur in parenchyma cells, provided that they do not contain calcium oxalate, either in solution in the cell sap, when the tissue is very young, or in a solid state in older parts. They are first formed in the leaves, and ultimately transferred to the bark.

On the other hand, caffeine and theobromine, which strictly speaking are purines, are generally considered to be decomposition products of proteins, ‡ they are formed in places of great cellular activity and their disappearance is never accompanied by a concomitant increase of albuminous substances.

These particular substances may correspond to urea and uric acid of higher animals, for the purine nucleus is characteristic of xanthine bases, such as uric acid; and derivatives of xanthine, such as guanine and adenine, are found in caffeine and theobromine. In this connexion one important point of distinction between animals and plants may be men-

^{*} Clautriau : loc. cit.

⁺ Lotsy: "Bull. Inst. Bot. Buitenzorg," No. 3, 1900.

[#] Clautriau: loc. cit.

tioned; in the higher animals there is a definite elimination of these waste nitrogenous substances from the organism, and the output bears a definite relation to the amount of proteins taken as food. In plants, on the other hand, there is no general elimination of nitrogenous waste, such substances being used up in anabolic processes. Thus Weevers,* whilst recognizing that caffeine and theobromine may be the products of the decomposition of proteins, considers that they are reorganized, and are therefore not to be classed as waste products in the same sense as uric acid is. It will, of course, be noticed that there is relatively much more nitrogen in these compounds than in the proteins.

Finally, some of the substances in question may be of biological importance as a protection against herbivorous animals and parasitic fungi.

No suggestions of any real value other than those already mentioned have as yet been made concerning the source from which these substances are synthesized, although, as pointed out by Meldola,† the discovery that glucose could by the action of ammonia in the presence of zinc hydroxide be converted into methyliminazole,‡ renders the genesis of some of the natural alkaloids which contain the iminazole ring at any rate a chemical possibility.

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*Weevers: "Proc. Koningkl. Akad. Wetens.," Amsterdam, 1903, 369; "Ann. Jard. Bot. Buitenzorg," 1907, 21, 1.
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SECTION VII.

COLLOIDS.

THE translocation of dissolved substances from cell to cell in the living plant is largely dependent on the diffusibility of liquids through colloidal membranes, and since some of the fluids occurring in plants are in reality colloidal solutions, it is desirable to consider briefly some of the more important properties of such solutions.

While studying the phenomenon of diffusion, Graham, in 1861, found that not all substances behaved in the same way when separated from water by a membrane of parchment or some similar material; thus, while such a substance as cane sugar passed through the membrane readily, albumen did not.

Crystallizable substances which, like cane sugar, were able to diffuse, he termed crystalloids, whilst those substances which, like starch, albumen, and gums, were not crystalline, and would not diffuse, he called colloids.

This difference may be illustrated by placing a mixture of starch paste and a solution of common salt in a parchment dialyser,* and floating this upon a large volume of distilled water. If a small portion of the mixture, and of the surrounding water, be tested from time to time with silver nitrate and with iodine solution, it will be found that the salt passes through the membrane, but the starch does not.

Graham further found that it was possible, under certain conditions, to cause otherwise insoluble substances to form colloidal solutions. Thus by adding an excess of dilute hydrochloric acid to a dilute solution of sodium silicate he obtained a clear solution instead of a precipitate of silicic acid. On subjecting this solution to dialysis, the sodium chloride was washed out, and there remained behind a clear liquid contain-

^{*} The dialyser may consist of a parchment bag, or of a tray formed by stretching a sheet of parchment over a circular wooden frame.

ing the ordinarily insoluble silicic acid in colloidal solution; this solution, which he termed a hydrosol, was stable at first, but after a few days it deposited silicic acid; the precipitated form of the colloid he termed a hydrogel, and since both dissolved and precipitated forms of colloid exist for many different solvents, the terms hydrosol, hydrogel, alcosol, alcogel, etc., were used to indicate the nature of the solvent.

The great difference in diffusibility between crystalloids and colloids is one of the most striking differences between these two classes of substances; it must not, however, be concluded that colloids are absolutely indiffusible, but rather that their rate of diffusion is very much slower than that of crystalloids.

Moreover, although a substance may, under given conditions, produce a colloidal solution, it does not follow that it will do so under all conditions; thus, a solution of soap in water is colloidal whilst a solution in alcohol is not; again, many substances, such as hæmoglobin, egg albumen, etc., which were formerly regarded as typical colloids, have been obtained in a crystalline form, while, on the other hand, most crystalloids can be made to give colloidal solutions; for instance, sodium chloride, if generated by some reaction taking place in benzene, gives a colloidal solution in that solvent.

Appended are some of the more important general characteristics of colloids.

GENERAL PROPERTIES OF COLLOIDS.

A. Diffusibility.

Graham considered that colloids were either incapable of diffusing through parchment or else possessed the power to a very limited degree; it would, however, appear that the diffusibility of colloids depends to some extent on the membrane, for a colloidal solution of congo red will diffuse through some varieties of parchment but not through others. In any case, however, the rate of diffusion is much slower than that of a crystalloid would be.

Osmotic Pressure.—Experimental determinations have shown that substances in colloidal solution produce a very small lowering of the freezing point or elevation of the boiling

opint of the solvent in which they are dissolved. From these observations it must be concluded either that the substances concerned have a very high molecular weight, or else that they exert a very small osmotic pressure,

Attempts to determine the molecular weights of substances by these means have given widely different and inconcordant results, the figures for egg albumen varying roughly from 6,000 to 15,000 according to different authors; moreover, while the depression produced by gallic acid in water corresponds to a very high molecular weight, similar measurements in glacial acetic acid give a molecular weight corresponding to that calculated from the formula. The conclusion to be drawn from these observations is that gallic acid gives a colloidal solution in water but not in glacial acetic acid.

In view of these facts the question arises whether the small depressions observed in the case of colloidal solutions are not really due to traces of impurities, such as salts, which it is almost impossible to remove completely.

On the other hand, the low osmotic pressure may be accounted for by assuming that colloidal solutions are in reality not true solutions, but are what may be described as two-phase systems; this implies that they are more of the nature of suspensions of one form of matter in another, such, for example, as a solid in a liquid; the suspended substance is known as the disperse phase, and the medium in which it is suspended as the continuous phase, a system of nomenclature which indicates that the particles of the liquid are in contact with each other, while those of the disperse phase are separate.

The above views are supported by much that is known regarding the optical properties of colloidal solutions.

B. Optical Properties.

The optical properties of colloidal solutions vary with the nature of the colloid; thus, for example, while a colloidal solution of silicic acid appears, to the unaided eye, homogeneous and clear, an aqueous solution of soap is more or less opalescent and polarizes light; on the other hand, while the colloidal solutions of organic substances are usually colourless, those of

the metals are frequently highly coloured; thus, gold may give a pink, red, or purple colour in water, and a blue colour in organic solvents, and, similarly, silver yields brownish-red or blue solutions.* This difference in colour given by the same metal depends upon the conditions under which the solutions are formed, and on the consequent size of the particles.

These facts, more especially those relating to opalescence and the polarization of light, indicate that such solutions are heterogeneous, i.e. they are not true solutions, but are more of the nature of suspensions of very minute particles. This conclusion is confirmed by the evidence afforded by the ultramicroscope, by the use of which the liquid can be seen to contain particles exhibiting different colours and Brownian movements. The size of the particles is very minute and varies greatly; some are visible under the high power of the ordinary microscope, whilst others are only just discernible under the ultramicroscope.

This may explain the fact that the particles of some colloidal solutions are able to pass through the pores of an ordinary filter paper, whereas they are stopped by the smaller pores of a parchment membrane; consequently, if the pores be relatively large slow diffusion could take place, but if the pores be so small as to allow only the molecules of a crystalloid, say of sodium chloride, to pass through, then the diffusion of colloids, e.g. starch paste, would be inhibited since the size of the particles is much greater than that of the molecules of salt.

It must, however, be remarked that such a passage of substances through membranes is understood imperfectly, and that many of the observed facts are incompatible with the filter theory expressed above.

The solution theory maintains that the membrane dissolves the diffusing substance which passes through and is given off on the other side.

The chemical theory holds that the passage of a substance through a membrane is due to the latter playing an active chemical part in the process.

^{*} Red, or so-called ruby, glass is also a colloidal solution of metallic gold in glass, and the naturally occurring blue rock salt is probably a colloidal solution of sodium in the crystalloid sodium chloride.

For a fuller account of these phenomena works on physical chemistry should be consulted.

C. Change of State of Colloids or Gel Formation.

It will be seen from what has gone before that, in addition to the normal colloidal solutions produced by dissolving typical colloids such as glue, gelatine, or albumen in water, there are a number of artificial colloidal solutions, some of which are produced from normally insoluble substances, such as silicic acid, gold, silver, and other inorganic substances, and some of which are produced from crystalloids such as sodium chloride.

With such widely different solutions it is not surprising to find a very considerable difference in stability, and, as a matter of fact, it is found that all colloidal solutions undergo with greater or less readiness a change of state resulting in their precipitation in the form of a gel. The resulting gel may be one which cannot be got into solution again, in which case the change is said to be irreversible; if, on the other hand, the change of state is only temporary, so that a slight change of conditions causes the re-solution of the gel, the change is said to be reversible.

The various types of reversible or irreversible change of state which colloidal solutions may undergo can be classified as follows:—

- (a) Spontaneous Precipitation, (b) Gelatinization, (c) Heat Coagulation, (d) Coagulation by Enzymes, (e) Precipitation by Electrolytes, (f) Precipitation by other Colloids.
- (a) Spontaneous Precipitation.—The decomposition of a colloidal solution of silicic acid after keeping for a few days may be quoted as an illustration of spontaneous precipitation.
- (b) Gelatinization by Altering the Concentration.—If a dilute solution of gelatine in water be concentrated until it is about 5 per cent strength it will set to a jelly on cooling to the atmospheric temperature. Solutions of agar will gelatinize even more readily. The change is, in both cases, reversible, for by raising the temperature, or by adding more water, the gel goes into solution again.
- (c) Heat Coagulation.—This change, which may be illustrated by the coagulation of egg white in boiling water, is irreversible.

An instructive experiment, due to Hardy, consists of boiling side by side in separate beakers a fairly strong and a very dilute solution of egg white in water. The strong one coagulates while the dilute one becomes turbid only; on the addition of a small quantity of barium chloride, however, a precipitate is produced. The explanation of this phenomenon is that owing to the dilution of the solution, the particles of coagulated protein are too small to unite together, and therefore remain apart forming a suspensoid which is, however, precipitated by the electrolyte.

(d) Coagulation by Enzymes.—The curdling of milk by rennet is a familiar example of this type of irreversible gel formation; so also is the coagulation of pectic bodies occurring in fruit juices by the enzyme pectase with the formation of gelatinous calcium pectate.

Enzymes capable of coagulating milk also occur in many plants, such as Lolium perenne, Anthriscus vulgaris, Geranium molle, Ranunculus bulbosus, Medicago lupulina, Ricinus, Datura, Pisum, Lupinus, etc.

(e) Precipitation by Electrolytes.—The effect of electrolytes on colloidal solutions differs very markedly according to the nature of the solution, since the inorganic colloidal solutions, or suspensoids, are extremely sensitive to the addition of only a small amount of an electrolyte, whilst, on the other hand, the so-called emulsoids, or organic colloidal solutions, are only affected by considerable quantities.

This is due to the fact that emulsoids have less well-defined electrical characteristics as compared with suspensoids (see next section on Electrical Properties).

The electrolytes which precipitate colloids from solution may be classified under two heads:—

- (i) Sodium, potassium, lithium, ammonium, and magnesium salts.
- (ii) Calcium, barium, strontium salts, and salts of the heavy metals, such as mercury, copper, lead, zinc, etc.

The former class produce reversible precipitation, while the precipitation produced by the second class is, as a rule, irreversible. Thus, for example, on saturating a solution of soap or gelatine, with ordinary salt, the colloid is precipitated in a form which can be redissolved in water at will. Practical application has been made of this phenomenon for separating the various types of protein. Thus, for example, if an aqueous solution containing an albumen and a globulin be mixed with an equal volume of saturated ammonium sulphate solution, the globulin, being insoluble in the resulting half-saturated ammonium sulphate, is precipitated; after filtering off the globulin, the albumen may be precipitated from the mother liquor by saturating it with ammonium sulphate.

The precipitated albumen and globulin are chemically unchanged and can be redissolved if desired.

Pauli,* in studying the precipitation of albumen by various salts, came to the conclusion that the precipitating power of a salt was an additive property which depended on the constituent ions.

Kations, as a rule, act as precipitants for albumen, while anions tend to keep it in solution.†

The precipitating power of the kations increases in the following order: Mg, NH_4 -, K, Na, Li, while the inhibiting or solvent action of the anion increases in the following order: $-C_2H_4O_2$ -,-Cl,-NO₃-,Br,-I,-CNS.

According as the precipitating power of the kation or the inhibiting power of the anion predominates the resulting salt will either precipitate or not precipitate albumen.

The observations are given below in tabular form. As shown by the arrows, the kations and the anions are arranged in ascending order of precipitating and inhibiting power respectively. The symbols + and - respectively signify that the salt does or does not precipitate albumen, the blank spaces meaning that the salt has not been investigated.

Kations	Mg	NH_4	K	Na	Li
Anions					
. ↓					
Fluoride		+	+	+	
Sulphate	+	+	+	+	+
Phosphate			+	+	+
Citrate			+	+	+
Tartrate			+	+	+
Acetate		-	-	+	+
Chloride	_	-	+	+	+
Nitrate	-	_	-	+	+
Chlorate			_	-	+
Bromide	-	-	-	-	+
Iodide	-	-			
Sulphocyanide	_	-	-	-	

^{*} Pauli: "Beitr. z. chem. Phys. u. Path.," 1902, 3, 225; 1903, 5, 30. + See p. 310.

From this table it may be seen that the comparatively slight precipitating power of the kations, Mg and NH₄-, is completely neutralized by the anions $-C_2H_3O_2$ or -Cl, while the more powerfully inhibiting anions -NO₃ and -ClO₃ are able to neutralize the precipitating power of the kation K as well as that of Mg and NH₄-. Similarly the powerfully inhibiting anions -Br, -I and -CNS, are able to counteract the precipitating power of sodium as well.

The salts of the metals belonging to the second class produce irreversible precipitation, owing, no doubt, to the formation of new compounds by chemical reactions. The case of zinc is peculiar, inasmuch as very dilute solutions of zinc salts produce irreversible precipitation of egg albumen, whereas strong solutions may either not produce a precipitate, or else cause one already formed to dissolve.*

The precipitating power of the anions when combined with one of the metals of the alkaline earths is exactly the reverse of that observed when the same anions were combined with the alkali metals. Thus the precipitating power of the anions increases in the order $C_2H_3O_2>Cl>NO_3>Br>I>CNS$, whereas when combined with the alkali metals the inhibiting power increased in this same order.

Irreversible precipitation of proteins is also brought about by nitric acid and by the alkaloidal reagents, such as picric, tannic, phosphomolybdic, phosphotungstic acids, etc.

(f) The Precipitation of Colloids by Other Colloids of Opposite Electric Sign.—This phenomenon was first observed by Linder and Picton, who found that certain solutions of organic dyes, on mixing, produced precipitates. Further investigations have shown conclusively that only oppositely charged colloids could mutually precipitate; thus, arsenic sulphide, which is negatively charged, is not precipitated by any other negatively charged colloid, but is precipitated by ferric hydroxide, which is positive. The resulting gel is described as an adsorption compound (see below under Adsorption).

This mutual precipitation of colloids has many very important practical applications; for example, the use of ferric salts in the purification of sewage water is probably due to the

^{*} Pauli: "Beitr. z. chem. Phys. u. Path.," 1905, 6, 233, 259.

precipitation of negatively charged colloidal particles of sewage by the ferric hydroxide hydrosol.

Similarly it has been suggested that the process of dyeing is really a mutual gel formation between the colloidal dye and the colloidal fibre; similarly the interaction between toxin and antitoxin, and the phenomenon of bacterial agglutination, etc., may be regarded as examples of the mutual precipitation of two colloids.

D. Protective Power.

Many organic substances such as gelatine, agar, etc., when added in small quantity to inorganic colloidal solutions, can prevent the precipitation of the latter by electrolytes; under these conditions the organic colloids are said to exert a protective action upon the inorganic colloid.

It is not known in what way this protective action is exerted, but it has been suggested that the particles of the suspensoid become covered with a layer of gelatine and so acquire the properties of gelatine particles.

Suspensoids, so protected, can be evaporated to dryness, and the residue when taken up with water will redissolve.

The greatly increased stability thus acquired by the inorganic colloid makes the process of value for the preparation of colloidal solutions of the metals, particularly silver and mercury, which are used for various medicinal purposes.

A measure of protective power was first worked out by Zsigmondy,* who defined as the gold number, the number of milligrams of colloid which, when added to 10 c.c. of a bright red colloidal gold solution containing from '0053 to '0058 per cent of gold, is just insufficient to prevent the precipitation (as shown by the colour change to violet) of the gold by I c.c. of a solution of sodium chloride, containing 100 grams of salt in 900 c.c. of water.

CLASSIFICATION OF COLLOIDS.

One of the most striking differences between organic and inorganic colloidal solutions is the much greater viscosity of the former. Thus, while even dilute solutions of gelatine are more or less sticky or viscous, inorganic colloidal solutions are

^{*} Zsigmondy: "Zeit. f. anal. Chem.," 1901, 40, 697.

never viscous. This difference is accounted for by assuming that the non-viscous solutions are, in reality, of the nature of suspensions of extremely minute solid particles in a liquid medium, whereas the viscous solutions consist of liquid particles suspended in a liquid medium.

Or, expressing it somewhat differently, it may be said that in the one case the disperse phase is a solid and the continuous phase a liquid, while in the other case both phases are liquid.

On this assumption rests the classification of colloidal solutions into Suspensoids and Emulsoids.

PROPERTIES OF SUSPENSOIDS

Although only the inorganic colloidal solutions belong to this class, so that they are relatively unimportant biologically, a brief description of their properties is essential to a survey of the whole subject.

The methods of preparation of inorganic suspensoids such as

- (a) The disintegration of metals by means of electric sparks under water;
- (b) The reduction of dilute solutions of salts of the metals by various reducing agents;
- (c) The passing of sulphuretted hydrogen through arsenious acid, whereby the arsenic trisulphide, instead of being precipitated, remains in colloidal solution,

all point to the presence of solid particles in such solutions.

Examination under the ultramicroscope reveals the fact that all these solutions contain extremely minute solid particles, the surface of which, from mathematical considerations, must be enormous in comparison to their mass. It is natural that such particles should be electrically charged owing to contact electrification, and they will therefore repel each other and so prevent aggregation to larger particles with consequent precipitation. This, moreover, accounts for their susceptibility to electrolytes, since the negatively charged particles become discharged by the positive ion of the electrolyte, whereupon the discharged particles are able to unite and form larger ones and so be precipitated.

This effect may be seen by the addition of a few drops of hydrochloric acid to a colloidal arsenic sulphide solution

when immediate precipitation results. Further, the assumption accounts for the fact that suspensoid colloids cannot be kept for any length of time unless carefully freed from electrolytes by dialysis.

The behaviour of suspensoids under the influence of a powerful electric current bears out the above views. In most cases the particles appear to be negatively charged since they travel towards the anode; exceptions to this are the hydrosols of the metallic hydroxides, such as iron, aluminium, etc., also silicic acid and basic dyes, such as methylene-blue and methylviolet, all of which are positive. It should, however, be noted that these charges are reversed if the same substances are dissolved in turpentine.

A fundamental difference between suspensoid colloidal solutions and true suspensions is that the latter when left at rest will ultimately deposit their suspended matter in the form of sediment as a result of gravitational attraction; by stirring up the sediment and again allowing it to settle the process can be repeated indefinitely, and it is, in fact, a reversible one, even though the sediment may have been heated or otherwise treated with a view to bringing about coagulation. A colloidal solution, on the other hand, is not in the same way affected by the force of gravity, and if effectually coagulated the change brought about is irreversible, and the precipitated substance will not go back into solution.

PROPERTIES OF EMULSOIDS.

Examination under the ultramicroscope reveals no distinct particles, but whether this is due to the particles being too small or not having a refractive index sufficiently distinct from that of the liquid in which they are suspended, is not as yet known.

As already stated, emulsoids are supposed to be composed of liquid particles consisting of a more concentrated solution suspended in a liquid medium composed of a much diluter solution. This assumption readily accounts for the viscosity of such solutions, which, being in reality suspensions of one liquid in another, would naturally be expected to have some of the properties of emulsions; that emulsions have a high

viscosity is well known from such a familiar example as codliver oil emulsion.

Furthermore, such a system of liquid particles with no sharply defined surface of separation from the surrounding medium would not be expected to exhibit any marked electrical properties due to contact electrification, such as occur in the case of solid particles. This accounts for their want of response to an electric current, and their relative indifference, as compared with suspensoids, to electrolytes when present in small quantities.

Whereas traces of acids or alkalis will at once precipitate suspensoids, similar quantities added to emulsoids will only have the effect of imparting to them slight electric properties in which they were previously deficient.

Thus, it has been shown by Hardy that whereas native albumen, when free from electrolytes, is electrically neutral, it acquires a negative charge on the addition of a little alkali, and a positive charge on the addition of acid.

According to Pauli,* this accounts for the fact that positively charged metallic hydroxides are unable to precipitate electrically neutral albumen, but precipitate albumen which has become negatively charged by the addition of a little alkali; and similarly negatively charged colloids, such as phosphomolybdic or phosphotungstic acid or certain negative dyes, are only able to precipitate albumen after it has acquired a positive charge by the addition of acid.

The behaviour of emulsoids towards stronger solutions of electrolytes is dealt with under the precipitation by electrolytes.

THE NATURE OF GELS.

As already pointed out above, emulsoids are regarded as two-phase systems in which the disperse phase is a more concentrated solution, and the continuous phase a relatively dilute one. When such a solution gives a gel, the rôles of the two phases are assumed to be changed, and one then has a sort of net- or sponge-like structure of concentrated solution representing the continuous phase, whereas the disperse phase is represented by a dilute solution filling up the interstices.

^{*} Pauli: "Beitr. chem. Phys. u. Path.," 1906, 7, 531.

Evidence for the existence of some such structure in gels is obtained from microscopic examination; furthermore the existence of so-called elastic gels, such as can be obtained by cooling gelatine solutions containing only about 5 or 10 per cent of solid and 95 or 80 per cent of water, explains to some extent how it is possible to obtain a rigid structure from plants such as asparagus or spinach, etc., which contain about 90 per cent of water.

In such a honeycomb or sponge-like structure there is, of course, a very large internal surface over all of which the phenomenon of absorption can take place. In consequence of this such gels will tend to retain with considerable tenacity small quantities of foreign substances, which accounts for the difficulty experienced in attempting to rid colloids of the last traces of impurities.

ADSORPTION.

The phenomenon known as the occlusion of gases is an example of the absorption of gaseous matter by a solid surface; it is exhibited to some extent by glass and platinum, but far better by wood charcoal, owing to its large superficial area; on this fact depends the use of wood charcoal as a deodorant or for the absorption of the last traces of gas in the production of high vacua. It is not known in what way the absorption is effected, but the immediate effect is to produce a concentration of gaseous molecules at the surface of contact between the solid and the gas.

Similarly, it has been shown from thermodynamical considerations, that when a solid body is introduced into a solution, the dissolved substance will tend to accumulate at the surface of contact between the solution and the solid.

To all such cases of purely surface absorption the term Adsorption is generally applied.

The concentration of a dissolved substance upon the surface of a solid introduced into a solution may be illustrated by dipping a piece of filter paper into a dilute aqueous solution of congo red; after a short time the dye will have accumulated on the surface of the paper, leaving the solution much lighter in colour.

Moreover, since congo red is in colloidal solution and filter

paper behaves in many respects like a colloid, this experiment also illustrates the phenomenon of mutual adsorption by colloids which is the principle underlying most processes of dyeing and staining, and also enzyme actions and other processes taking place in the living organism.

It is, of course, easy to understand that if adsorption takes place so readily between colloids, such as filter paper and congo red, both of which bear negative charges in water, the phenomenon must take place still more easily between oppositely charged colloids in which the mutual electrical discharge facilitates the deposition.

Numerous practical applications of adsorption from solutions are known, as for example in the removal of colouring matter in the purification of cane sugar, or in the removal of fusel oil from crude spirit by filtration through charcoal.

Other substances besides charcoal, such as fuller's earth and china clay, have been similarly used on account of the large surfaces which they present.

From what has been said with regard to the structure of gels and the assumption that they present a sort of network with a considerable development of internal surface, it is easy to find an explanation of the use of isinglass for clearing a turbid solution or for the fact that colouring matter may be extracted from a solution by precipitating gelatinous aluminium hydroxide in it.

Thermodynamical considerations, coupled with experimental measurements, show the fact that true adsorption takes place according to well-defined mathematical laws which enable one to decide definitely whether a certain phenomenon is due to physical adsorption or to chemical reaction; thus, it has been found that a relatively larger amount of the total substance in solution is withdrawn from a dilute than from a strong solution.

It has been calculated that the total surface presented by the particles of a red colloidal gold solution containing 0.5 gram of gold per litre amounts to about 8 sq. metres. It is, therefore, easy to understand that with such an enormous development of surface there is the possibility for a marked manifestation of adsorption by suspensoids, and this readily explains the tenacity with which most such colloidal solutions retain traces of electrolytes, and why so many precipitates, in spite of continued washing, persistently retain small quantities of substances which they have carried down with them from solution.

The following experiment, due to Linder and Picton,* illustrates this phenomenon:—

If, to a colloidal solution of arsenic sulphide, obtained by passing a current of sulphuretted hydrogen through arsenious acid dissolved in water, a little barium chloride be added, the arsenic sulphide is precipitated. This precipitate, if filtered and washed thoroughly to free it from barium salts, will still be found to contain traces of barium if dissolved in concentrated hydrochloric acid with a little potassium chlorate and tested for barium in the usual way.

Again, the mutual precipitation of colloids of opposite electrical sign,† resulting in the formation of a coagulum consisting of both constituents, is probably only another example of adsorption.

Many natural phenomena can be attributed to the same cause. For example, the power possessed by soils rich in clay or humus to retain soluble potassium salts or phosphates which would otherwise be washed away by rain.

The hydrated aluminium magnesium and sodium silicates, known as Zaeolites, which are contained in clays are colloids and they react by double decomposition with the potassium salts which may be applied as manures, and, while retaining the potash, set free a corresponding quantity of lime or soda.

Another very striking case of selective adsorption is to be found in the power which sea-weeds have of extracting iodine from the surrounding sea-water, although the amount of this element in sea-water is extremely small; again, in spite of the enormous preponderance of sodium over all other metals in sea-water, the plant takes up practically none of this, but takes instead potassium, which is present in much smaller quantity.

ENZYME ACTION OF COLLOIDS.

Associated with this enormous development of surface there is, of course, a corresponding development of surface

^{*} Linder and Picton: "J. Chem. Soc., Lond.," 1895, **67**, 66. + Cf. van Bemmelen: "Z. anorg. Chem.," 1900, **23**, 300.

energy* which no doubt, in part, explains the remarkable catalytic activity exhibited by colloidal solutions of the metals.

Bredig† and his collaborators have shown that a colloidal solution of platinum containing 194 grams of metal (i.e. I gram atom) in 70,000,000 litres of water, or a colloidal solution of gold containing 197 grams of metal in 1,000,000 parts of water, are still able to produce a distinct accelerating influence on the decomposition of hydrogen peroxide into water and oxygen.

It has long since been known that metallic platinum, more especially the variety known as spongy platinum, when left in contact with hydrogen peroxide induces the decomposition of this substance into water and oxygen, and Berzelius,‡ as long ago as 1836, pointed out an analogy between this catalytic action of platinum and the action of an insoluble ferment, such as yeast on sugar.

This suggestion has since been borne out by a number of examples of chemical changes which could be effected equally well either by means of finely divided platinum or by a ferment, e.g. the oxidation of alcohol to acetic acid by *Mycoderma aceti*, the bleaching of indigo solution by hydrogen peroxide in presence of red blood corpuscles, the blueing of tincture of guaiacum by hydrogen peroxide in presence of red blood corpuscles, etc., all of which can also be effected by spongy platinum.

Bredig carried our knowledge of the subject a step farther; by preparing colloidal solutions of the metals and comparing their action with that of various enzymes, he traced out such a remarkable analogy between the two that he has called the colloidal metal solutions "Inorganic Ferments".

The chief points of similarity between enzymes and colloidal platinum may be summarized as follows:—

- Both platinum hydrosol and enzymes are colloids and as such are detrimentally affected by electrolytes.
- 2. Both platinum hydrosol and enzymes gradually decompose spontaneously or decompose more rapidly by heating.

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* Ostwald: "Z. physik. Chem.," 1897, 23, 172.
† Bredig: "Anorganische Fermente," Leipsig, 1901, p. 96.
‡ Berzelius: "Jahresber.," 1836, 13, 237.
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- 3. There is an optimum temperature for both colloidal platinum and for enzymes to exert their catalytic action.
- 4. The activity of platinum hydrosol may be stimulated by the addition of alkali until it reaches its maximum value, after which the further addition of alkali causes it to fall again. Similar stimulation of enzymes by the addition of certain substances known as Zymo-exciters have been observed in case of emulsin acting on hydrogen peroxide, and of invertase acting on cane sugar.
- 5. The decomposition of hydrogen peroxide whether by platinum hydrosol or by hæmase, the enzyme contained in blood, is in accordance with the laws governing a monomolecular reaction.
- 6. A very remarkable analogy between platinum hydrosol and the enzyme of blood is that small quantities of substances which, when added to the colloidal platinum solution, destroy its catalytic action on hydrogen peroxide, also have the same effect on the oxidase of blood. Curiously enough many of these substances are blood poisons such as sulphuretted hydrogen, hydrocyanic acid, carbon monoxide, and arseniuretted hydrogen; several other substances were also found to paralyse either the platinum solution or the enzyme.

It was further observed that platinum hydrosol when treated with very small traces of hydrocyanic acid was temporarily poisoned but recovered after a short time; a similar effect has also been observed with enzymes. The recovery is probably due to the oxidation of the hydrocyanic acid.

It was also found that the toxic effect of the hydrocyanic acid was much greater if added directly to the platinum or gold sol than if added to a sol already containing some hydrogen peroxide. Exactly similar conditions had been previously found by Schönbein* to hold in regard to the addition of hydrocyanic acid and hydrogen peroxide to blood.

In conclusion it should be noted that Bredig, while disclaiming any attempt to trace a fanciful connexion between the colloidal metal solutions and enzymes, emphasizes the fact that the two properties of catalytic action and colloidal nature are common to both classes of compounds and regards the

^{*} Schönbein: "Zeit. f. Biologie," 1867, 3, 144.

colloidal metals as the simple inorganic analogues of the more complex enzymes.

One further illustration might be quoted of the chemical activity which is associated with colloidal substances presenting a large surface. A calculation based on the assumption that there are five million red blood corpuscles of diameter '007 mm. contained in 1 c.mm. of blood reveals the striking fact that the total surface presented by the blood corpuscles contained in 5 litres of blood (the amount contained in the body of a full-grown man) would be about 1875 square meters. From what has gone before it is, therefore, not surprising that these corpuscles should be endowed with special properties enabling them, in the presence of the trace of iron which they contain, to play their part in the highly complex changes involved in respiration.

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SECTION VIII.

PROTEINS.

THE term protein is applied to a large variety of bodies occurring in the animal and vegetable kingdoms, which occupy a pre-eminent position in the economy of life, owing to their being the chief constituents of protoplasm.

In the plant, proteins may occur either as solid bodies or in solution in the cell sap. They may be found in all living members; in roots, stems, leaves, sieve tubes, laticiferous tissue, etc. Reserve proteins commonly are found in the solid state, especially in seeds and in vegetative organs of propagation.

These protein bodies may be either quite amorphous or crystalline; sometimes the grains are partly amorphous and partly crystalline, as in the well-known aleurone grains of the seed of *Ricinus* (castor oil).

Protein crystals may be cubical, as in the potato, falciform as in the carpellary walls of *Gratiola officinalis*, and other shapes; they may occur quite free within the cell, as in the potato, or embedded in other bodies. These embedded crystals may be found in nuclei, e.g. in the leaves of *Melampyrum arvense* and in the ovary wall of *Campanula trachelium*; in chloroplasts, e.g. *Hedera* and *Canna*; and in amorphous protein, e.g. in the seeds of *Ricinus* and *Berthelletia*.

These last, generally known as aleurone grains, are often somewhat complicated; the grain is surrounded by a protein membrane, which is less readily soluble than the remaining amorphous protein of the matrix. Embedded in the matrix is the crystalloid, and also a globoid consisting of a double phosphate of calcium and magnesium. The crystalloids vary in shape; commonly they are hexagonal and stain brown with iodine and are readily soluble in dilute alkali. Also they

may readily be stained in fuchsin. To do this, the sections should be placed in a '2 per cent aqueous solution of acid fuchsin for twenty-four hours, washed in running water and mounted in Canada balsam in the usual way.

Several proteins may occur in aleurone grains and may be recognized by their different solubilities in water, salt solution, alkali, and alcohol. Also, the details of the composition of these grains are not the same for all plants in which they occur; for instance, in the Pæony the matrix is soluble in water, whereas in the castor-oil plant it is insoluble in water, but soluble in a strong aqueous solution of sodium phosphate.

According to Bokorny,* globulins are the common proteins occurring in the aleurone grains and crystalloids of seeds. It should be remarked that the term aleurone grain is frequently used in a generic sense to include all non-crystalline reserve protein bodies of a more or less definite shape; they are not always of the complicated nature described above, thus in the grain of wheat they are quite simple in structure and do not contain a crystalloid nor a globoid.†

GENERAL PROPERTIES OF PROTEINS.

Until recently, comparatively little was known with regard to the chemical nature of proteins beyond the fact that they were composed of the elements carbon, hydrogen, nitrogen, oxygen and sulphur, together with, in some cases, phosphorus and iron; their existence as a separate group of compounds therefore depended chiefly on their sharing a number of general physical and chemical properties, without regard to their constitution, concerning which little or nothing was known.

A scientific definition of proteins was first given by Panzer, who classified as proteins all substances which on hydrolysis yield mono- or di-amino acids. This definition is, however, too comprehensive, as it would include amongst the proteins the group of substances described by Fischer as polypeptides.

The general physical and chemical properties which are shared by the typical unaltered proteins, such as albumins and globulins, may be summarized as follows:—

^{*} Bokorny: "Bot. Centrbl.," 1900, 82, 289.

⁺ For an account of the artificial production of protein grains, see Thompson: "Bot. Gaz.," 1912, 54, 336.

Panzer: "Wiener klin. Wochenschr.," 1903, 16, 689.

A. Physical Properties.

- Indiffusibility.
 Coagulation.
 Optical activity.
 - 4. Precipitation without change.
- B. CHEMICAL PROPERTIES. [Shared by all proteins.]
 - 1. Precipitation reactions. 2. Colour reactions.

A. Physical Properties.

1. Indiffusibility.

Unaltered or native proteins* belong to that class of bodies known as colloids (see p. 283) which are unable to diffuse through a parchment or animal membrane; it is thus frequently possible to purify a protein from salts by dialysis. The separation is, however, not quantitative, and it has, hitherto, not been found possible to remove from any protein the last traces of adhering inorganic salts, so that a perfectly pure protein, which on ignition yields no ash, has not as yet been obtained by this means.

Although all proteins are more or less colloidal in nature, they possess this property of diffusion in a varying degree; thus, for example, the albumoses and peptones, which are derived from the more complex proteins by the action of certain ferments, diffuse with comparative ease. In view, however, of the fact that these substances have never as yet been obtained in crystalline form, and that such typical colloids as oxyhæmoglobin and serumalbumin have been crystallized, the rigid distinction between colloid and crystalloid can no longer be upheld.

2. Coagulation.

All genuine or native proteins on keeping undergo a curious change known as coagulation; the nature of this change is at present not understood, but as a result of it, such proteins lose their distinctive properties of solubility, and can no longer be dissolved without first decomposing them into simpler substances, as, for example, the albumoses or peptones.

Coagulation may be brought about by (a) Heat, (b) Ferments, (c) Alcohol.

(a) The solutions of all albumins or globulins may be coagulated by heating; the temperature at which the change

^{*} The term native protein is applied to proteins which have been isolated from the tissues by some simple process which does not involve any alteration in their original properties.

takes place is characteristic for each substance, and varies from 56° C. in the case of fibrinogen to $70\text{-}80^{\circ}$ C. for serum albumin.* The reaction of the solution as well as the presence of dissolved salts are factors which exercise a powerful influence, a slightly acid solution being most favourable for the phenomenon, whereas an alkaline reaction may prevent coagulation entirely. According to Blum, formaldehyde is also able to prevent heat coagulation.

Heat coagulation is best effected as follows. The solution is first boiled, and from 1-3 drops of dilute acetic acid are added for each 10 c.c. of liquid, according to the amount of protein present, the liquid being boiled each time before the addition of each drop.

If the amount of salts present be small, a little I per cent sodium chloride should first be added, as the precipitation of small quantities of protein cannot otherwise be guaranteed.

- (b) Some proteins are rendered insoluble by the action of certain ferments, e.g., the precipitation of casein from milk by the action of rennet on caseinogen.
- (c) The addition of absolute alcohol to a neutral or faintly acid solution of a native protein will precipitate it from solution unchanged. If, however, it be left in contact with the alcohol for some time, the protein is rendered insoluble and is coagulated.

The solution must not be alkaline, and must contain a small quantity of neutral salts.

3. Optical activity.

The solutions of all proteins are lævo-rotatory, the amount varying from - 33.5° in the case of egg albumen to - 80° in the case of casein.

4. Precipitation without change.

Certain salts, such as sodium chloride and the sulphates of sodium, magnesium and ammonium, etc., have the property of throwing proteins, except peptones, out of solution. This is, however, purely a physical phenomenon, and must be distinguished from the chemical precipitation described below,

^{*} The coagulation temperature is not sufficiently well defined to be employed as a means of identification.

inasmuch as the proteins are precipitated unchanged, and retain all their original properties and solubilities. Absolute alcohol, also, as mentioned above, precipitates the proteins unchanged, though the precipitate must not be left in contact with the alcohol, or else it will become coagulated.

With regard to the precipitating power of these various salts, it should be mentioned that saturated ammonium sulphate will precipitate *all* proteins except peptones, and consequently a solution which on saturation with ammonium sulphate remains clear, can be regarded as free from protein.

Furthermore, zinc sulphate is approximately equivalent to saturated sodium chloride is approximately equivalent to

ammonium sulphate.

saturated magnesium sulphate, or 1/2 saturated ammonium sulphate.

In view of the number of proteins in the plant and their different characteristic solubilities, it is easy to see the importance to the well-being of the plant of factors which have a bearing on these properties. Thus any cause which removes water, not immediately replaceable, from the cell, and so leads to a concentration of the cell sap, may be a determining factor in the existence of a plant. Cold is one such factor;* a fall in the temperature may cause the water to crystallize, so that the salt solutions in the cell become stronger, with the result that some of the proteins of the protoplasm may be dissolved and other proteins in solution may be precipitated. The importance of soluble carbohydrates and of oils in the cell sap in this connexion has already been pointed out.

It is unnecessary to remark that this effect of cold must vary pretty considerably in different plants, and depends upon the nature of the salts dissolved in the cell sap and the proteins upon which they can act. To take a few examples: it was found that in Begonia, soluble proteins were precipitated when the temperature reached -3° C.; on the other hand, in the leaves of Pinus, a temperature of -40° C. was required to obtain a similar result.† This may, in part, be due to the paucity of crystalloids in the cell sap, for it is stated that

^{*} See Blackman: "New Phytol.," 1909, **8**, 354. + Gorke: "Landwirth. Versuchs. Stat.," 1906, **65**, 149.

plants which are subject to periodic drought possess only small amounts of soluble crystalloids in the cell sap.

In the case of the barley, it was observed that an exposure for one night to a temperature of -7° C. reduced the yield of soluble proteins by about one-third as compared with a control experiment in which the temperature was not so lowered. This salting out effect is much increased if the cell sap becomes acid on cooling, as is not infrequently the case.

If the low temperature be long continued, the precipitated proteins will not again enter into solution when the amount of water is increased by raising the temperature; on the other hand, if the temperature be suddenly raised, the precipitated proteins will re-dissolve, provided that they have not stood too long, and thus the plant will not be greatly harmed.

B. Chemical Properties.

1. Precipitation reactions.

The proteins have both acid and basic properties; thus, casein may be looked upon as typically acid, seeing that it dissolves in alkalis to form sodium and potassium salts, whilst the histones and protamines are powerful bases. All proteins, however, have basic properties, which enable them to form insoluble salts with a great many of the ordinary alkaloidal reagents, such as phosphotungstic, tannic, picric, ferrocyanic, and trichloro-acetic acids. They are also precipitated by potassium ter-iodide (a solution of iodine in potassium iodide) and by the double iodides of potassium with mercury, bismuth, and cadmium.

The strong mineral acids also precipitate proteins.

In consequence of this dual nature of proteins they are classed as amphoteric electrolytes (see below).

The salts of the heavy metals also produce insoluble precipitates with the proteins, a fact which is made use of in the administration of egg albumen as an antidote in cases of poison with salts of the metals. Moreover, the antiseptic action of mercuric chloride is most probably connected with this formation of insoluble salts.

Amongst the salts most frequently used as precipitants for proteins are the chloride and acetate of iron, the sulphate and acetate of copper, the chloride of mercury and the acetates of lead and zinc.

2. Colour reactions.

These reactions depend on the fact that certain groups or radicles in the protein molecule produce characteristic colours with suitable reagents. The reactions may also be employed for detecting these same groups in the decomposition products of the proteins, with the object of determining how far the decomposition has gone, and whether it has been sufficiently deep-seated to destroy this grouping or not. The following is a list of the more important colour reactions:-

(i) Biuret Reaction.—This is the bluish-violet colour produced by adding copper sulphate to an alkaline solution of a protein. Unchanged proteins give a bluish-red, whilst altered proteins, such as the peptones, give a pink.

The colour is given by the substance biuret itself, whose composition is expressed by the formula NH₀CO.NH. CONH₀, and by similarly constituted compounds containing two -CO. NH- groups connected together through a carbon, nitrogen, or sulphur atom.

- (ii) Millon's Reaction.—A solution of mercuric nitrate containing nitrous acid added to a solution of a protein produces a precipitate which turns pink or red. This reaction is connected with the phenolic group of the tyrosine complex in the protein molecule; it may also be used as a test for tyrosine.
- (iii) Xanthoproteic Reaction.—Protein solutions treated with concentrated nitric acid develop a yellow colour which is intensified by heating, and is changed to orange by ammonia. This reaction is likewise connected with the tyrosine complex.
- (iv) Adamkiewicz's Reaction.—The addition of concentrated sulphuric acid to a solution of a protein dissolved in acetic acid produces a reddish-green or violet colour. This reaction is characteristic of the tryptophane group, and is produced by the interaction of this with glyoxylic acid contained in the acetic acid; the reaction may be intensified by replacing acetic by glyoxylic acid in the test.
- (v) Liebermann's Reaction.—Proteins, which have been previously extracted with alcohol and ether to remove fats. on warming with concentrated hydrochloric acid, develop a

violet colour. According to Cole,* the colour is due to the presence of glyoxylic acid as an impurity in the ether.

(vi) Molisch's Reaction is a reaction for furfurol produced by the action of concentrated sulphuric acid on a carbohydrate. The substance to be tested is treated with a few drops of a 10 per cent alcoholic solution of α -naphthol; concentrated sulphuric acid is then slowly added, when a red-violet colour is formed at the junction of the two liquids.

Microchemical Reactions.

The following are the usual microchemical tests employed for the indication of proteins within the plant:—

- 1. Iodine gives a yellow to brown coloration.
- 2. With osmic acid a brown coloration results.
- 3. Biuret reaction.—A solution of copper hydrate in caustic potash may be added direct to the preparation; or the section may be steeped for some time, say twenty to sixty minutes, in 2 per cent solution of potash, washed, placed in a 10 per cent solution of copper sulphate for thirty to sixty minutes, washed in water and mounted in a 2 per cent solution of caustic potash. A mauve to violet coloration indicates the presence of proteins.
- 4. Millon's reagent.—The section or scraping is mounted in a few drops of the reagent and warmed. A brick-red coloration results when proteins are present. The reagent may be prepared by dissolving some mercury in twice its weight of nitric acid (sp. gr. 1·42), the operation being performed in a fume cupboard. When the action has ceased, the solution is diluted with twice its volume of water.
- 5. Xanthoproteic reaction.—A yellow to orange coloration results with proteins. The preparation is warmed on the slip with a few drops of strong nitric acid. The proteins acquire a yellow colour which is changed to orange on moistening with strong ammonia.

PROTEINS AS COLLOIDS.

An important difference between the solution of an electrolyte and of a colloid lies in the fact that whereas the electrolyte breaks up into two oppositely charged ions, the colloid appears

^{*} Cole: "Journ. Physiol.," 1904, 30, 311.

to be charged as a whole either positively or negatively, and accordingly when such a solution is subjected to a difference of potential, the colloidal particles wander bodily to one or other of the electrodes.

Biltz has shown that two colloids mutually precipitate each other only if they bear unlike charges, and when once precipitated, they become electrically neutral and are no longer transported by an electric current. It has further been shown that only those crystalloids which are electrolytes are able to precipitate colloids, such substances as urea or cane sugar being unable to effect precipitation.

According to Hardy and Bredig, the particles in a colloidal solution are held in suspension by the opposing forces of capillary attraction and of electrostatic repulsion such as must exist between particles all of which bear the same charge; the precipitation of a colloid by the addition of an electrolyte is accordingly attributed to the elimination of the electrostatic repulsion by the fact of the charge borne by the ions of the electrolyte neutralizing the charge borne by the particles. Billitzer, on the other hand, holds the view that on introducing the electrolyte into the solution the particles tend to congregate around the ions, and are thereby brought into such close contact with each other that they form sufficiently large aggregates to be precipitated.

By means of conductivity experiments, Pauli* was able to show that pure egg albumen, free from electrolytes by repeated dialysis, was electrically neutral, for, on subjecting a solution of this substance to an electric current for twenty-four hours, no particles of albumen were transferred to either of the two electrodes. He found, moreover, that the addition of neutral salts of the alkali metals or the metals of the alkaline earths, produced no change in the electrical state of the albumen, whereas on adding traces of acids or acid salts the albumen assumed a positive charge, while on the addition of bases or salts having an alkaline reaction, it became electronegative. This observation that an electrolyte, such as hydrochloric acid, which contains an equal number of oppositely charged ions, is able to impart a positive charge to electric-

^{*} Pauli: "Hofmeister's Beiträge," 1902, III, 225; 1904, V, 27; 1905, VI, 233, etc.

ally neutral albumen, is explained by assuming that albumen exerts some selective action on the ions, and is more permeable to positively charged hydrogen ions than it is to the negative chlorine ions. Pauli has further shown that his electrically neutral albumen, unlike ordinary albumen, is not precipitated from solution by the addition of salts of copper, iron, zinc, lead or mercury. The fact that neutral albumen is, however, precipitated by alcohol, although both substances are electrically neutral, must not be taken as evidence against the view that precipitation is produced as a result of the neutralization of the charges borne by colloidal particles; the explanation of the precipitation in this case lies in the complete insolubility of albumen in alcohol.

These facts throw some light on the electrical behaviour of native albumen in the living organism, for inasmuch as salts of the metals at once precipitate such albumen, whereas they refuse to precipitate neutral albumen, it follows that native albumen must bear a negative charge. This negative charge is most probably produced by the hydroxyl ions * liberated from the salts in contact with it, a view which receives support from the fact that on adding sodium bicarbonate to a fresh solution of neutral albumen, the latter at once assumes a negative charge.

As a consequence of this negative charge, it follows that the greater the electro-positive nature of an element, the greater will be its tendency to precipitate native albumen; on the other hand, the electro-negative acid radicles will tend to prevent precipitation, a tendency which is found to increase in the following order—sulphate acetate, chloride, nitrate, bromide, iodide and sulphocyanide.

The antagonistic action of electro-positive and electronegative radicles accounts for the fact that sodium in the form of sodium sulphate is a precipitant, whereas in the form of sodium bromide it is not; in the case of sodium iodide and sulphocyanide, the influence of the electro-negative radicle entirely outweighs that of the electro-positive sodium, with the result that these salts not merely do not precipitate albumen themselves, but actually interfere with the precipitation of albumen by other salts.

^{*} Pauli: "Hofmeister's Beiträge," 1906, 7, 531.

			-SO ₄	-C ₂ H ₃ O ₂	—C1	-NO ₃	—Br	-1	-cns				
Li	•		+		+	+	+						
Na			+	+	+	+	_						
К			+	+	+		-	_					
NH					_	_	-	_	_				

Increasing Tendency to Prevent Precipitation of Electro-Negative Albumen.

The sign + indicates that the reagent precipitates albumen, and - that it does not.

If, however, we make an albumen solution electro-positive by the addition of a little acid, the electro-negative or acid radicles now become the precipitants, and the whole order is reversed, and those salts which like the bromide, iodide or sulphocyanide, tended to prevent precipitation, now become the most powerful precipitants.

AMPHOTERIC ELECTROLYTES.

Amphoteric electrolytes are defined by Bredig* as substances which in aqueous solution are able to exhibit both acid and basic properties, and are accordingly able to split off or combine with -H or -OH ions.

Thus, in the presence of bases they behave as acids and dissociate as follows:—

$$ROH = RO + H$$

whereas in the presence of acids they behave like bases, giving the following ions:—

$$ROH \rightleftharpoons R + OH$$

This phenomenon of changing electrolytic distribution according to circumstances is known as "electrolytic tautomerism". Examples of amphoteric electrolytes in inorganic compounds are to be found amongst the hydroxyl derivatives of most of the elements from the middle of the periodic table,

^{*} Bredig: "Zeit. Electrochem.," 1899, 33.

such as aluminium, chromium, zinc, lead, tin, manganese, arsenic or antimony, all of which are weak bases or acids.

The fact that egg albumen is able to neutralize hydrochloric acid was first observed by Sjöqvist,* and later by Bugarsky and Liebermann;† moreover, the fact that both basic dyes such as rosanilines, as well as acid dyes such as picric acid, are able to combine with wool fibres, points to the amphoteric nature of the protein in this case also.

The simultaneous possession by a body of both acidic and basic groupings; is well illustrated by the amino acids; the true amphoteric character is, however, only illustrated by those of them that are sufficiently weak acids, or whose acidic and basic functions are about equally strong.

The observation that the hydrochloride of albumen is precipitated by the addition of sodium phosphomolybdate, whereas albumen itself is not, led Spiro and Pemsel § to conclude that albumen belongs to a class of compounds which, although electrically charged, are not ionized, and while not functioning as acids or bases themselves, are none the less able to form addition compounds with such substances. According to Sjöqvist | and Bugarsky and Liebermann, albumen forms with acids and bases true salts, which obey all the laws of Van't Hoff and Arrhenius, though, on the other hand, its low conductivity would appear to preclude the possibility of its possessing well-marked hydrogen or hydroxyl ions. These facts are, however, readily explained by assuming proteins to be pseudo-bases of the type described by Hantzsch** and his collaborators. Hantzsch describes as pseudo-bases a class of compounds which are chemically indifferent, but which on coming in contact with acids undergo molecular rearrangement, giving rise to true bases.

This change, which may be represented as follows,

 $N \equiv ROH \rightarrow R \equiv N \cdot OH$

implies a change from a neutral carbinol to a true ammonium

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* Sjöqvist: "Skand. Arch. Physiol.," 1895, 5, 354.
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⁺ Bugarsky and Liebermann: Pflüger's "Archiv Physiol.," 1808, 72, 68.

[‡]Winkelblech: "Zeit. physikal Chem.," 1901, 36, 551.

[§] Spiro and Pemsel: "Zeit. physiol. Chem.," 1898, 26, 270. || Sjöqvist: "Skand. Arch. Physiol.," 1895, 5, 277.

[¶] Bugarsky and Liebermann: "Zeit. gesam. Physiol.," 1898, 72, 68.
** Hantzsch: "Ber. deut. chem. Gesells.," 1899, 32, 575, 3100; 1900, 33, 278.

base with pentavalent nitrogen, which is able to react with acids to form salts of the type of $R \equiv N \cdot Cl$.

This assumption explains the reason why albumen is not precipitated by sodium phosphomolybdate until a little acid has been added, for if albumen in neutral solution is a pseudo-base, it would only be converted into a true base capable of being precipitated by sodium phosphomolybdate after the addition of acid.

For a further account of the connexion between pseudo-acids or bases and amphoteric electrolytes, see Zadwidzki* and Hantzsch.†

THE DECOMPOSITION PRODUCTS OF THE PROTEINS.

The most direct way of obtaining an insight into the probable groups or groupings which occur in the molecule of some complex substance, is to break it up into simpler ones, whose constitution is already known, or may be determined with comparative ease. This is the method which has been employed to elucidate the very complex structure of the proteins

Various processes have been employed for breaking down the protein molecule, such as acid hydrolysis, fusion with alkalis, the action of enzymes or putrefactive bacteria, oxidation, etc. As a result of all these various methods, a number of simple compounds have been obtained, which fall primarily into two main groups:—

- I. *Biuretic derivatives*, such as albumoses, peptones, etc., which are still very complex substances, but have, at any rate, a lower molecular weight than the original unaltered protein. These substances all give the Biuret reaction.
- 2. Abiuretic derivatives.—In this group of cleavage products, which give no Biuret reaction, are included the various amino acids.

By an amino acid is meant an acid in which one or more of the hydrogen atoms other than the carboxylic hydrogen are replaced by the amino group —NH₂. Thus acetic acid CH₃COOH gives rise to the amino acid known as glycine CH₃NH₃COOH. Theoretically it should be possible to re-

^{*} Zadwidzki: "Ber. deut. chem. Gesells.," 1903, **36**, 33²5; 1904, **37**, 153. † Hantzsch: "Ber. deut. chem. Gesells.," 1906, **39**, 3149.

place two or even three atoms of hydrogen in acetic acid by the $-NH_2$ group to produce diamino acetic acid $CH(NH_2)_2COOH$ and triamino acetic acid $C(NH_2)_3COOH$; these compounds are, however, not known, and appear to be incapable of existing.

The next homologue after acetic acid, namely, propionic acid CH_2CH_2COOH , can give two mono-amino acids CH_3CHNH_2COOH and $CH_2NH_2CH_2COOH$ known respectively as a- and β - amino propionic acids, according as the amino group is attached to the a- carbon atom, adjacent to the carboxyl group, or to the β - carbon atom, which is next but one from the carboxyl.

In the case of the higher homologues, diamino acids are known which have two amino groups attached to different carbon atoms, such, for example, as α-δ-diamino valeric acid CH₂NH₂CH₂CH₂CHNH₂COOH derived from valeric acid CH₃CH₃CH₃CH₃COOH.

The dicarboxylic acids also can give rise to amino derivatives such as aspartic acid COOHCH₂CHNH₂COOH derived from the dicarboxylic acid succinic acid COOHCH₂CH₂COOH and glutamic acid COOHCH₂CH₂CHNH₂COOH derived from glutaric acid COOHCH₃CH₃CH₂COOH.

It is important to note that all amino acids which are known to take part in the building up of the protein molecule are asubstituted acids, as will be seen from the list of protein cleavage products given below.

The presence of the —NH₂ group in amino acids confers upon these substances basic properties, in addition to the acid properties which they already possess. Thus, for example, glycine CH₂NH₂COOH is able to react with hydrochloric acid to produce glycine hydrochloride CH₂NH₂HClCOOH, just as ammonia reacts with hydrochloric acid to give a hydrochloride; on the other hand, being an acid, it is also able to form metallic salts, such as CH₂NH₂COOK. It is not surprising to learn that the mono-amino acids, such as glycine and its homologues, have no very pronounced acidic or basic properties; they belong, in fact, to the class of bodies known as amphoteric electrolytes (see p. 311). On the other hand, the mono-amino derivatives of the dicarboxylic acids, namely, aspartic acid COOHCH₂CHNH₂COOH and glutamic acid COOHCH₂

CH₂CHNH₂COOH, are strong acids, owing to the predominating influence of the two carboxyl groups, while the diamino derivatives of the monocarboxylic acids, such as lysine CH₂NH₂CH₂CH₂CH₂CHNH₂COOH, ornithine CH₂NH₂CH₂CH₂CH₂CHNH₂COOH, etc., have strongly marked basic characteristics, the two amino groups here overpowering the single carboxyl group.

A class of substances which have to be carefully distinguished from the amino acids are the *acid amides*. These are derived from carboxylic acids by replacing the hydroxyl group of the carboxyl by —NH₂. Thus acetic acid CH₃COOH gives the amide CH₃CONH₂ known as acetamide, while aspartic acid COOHCH₂CHNH₂COOH gives the amide CONH₂CH₃CHNH₂COOH known as asparagine.

AMINO ACIDS OBTAINED AS CLEAVAGE PRODUCTS OF

(I) Aliphatic Compounds.

(Glycine or a-amino-acetic acid CH₂NH₃COOH Alanine or a-amino-propionic acid CH2CHNH2COOH Valine or α-amino-isovaleric acid CH₃ CH . CHNH₂COOH Leucine or a-amino-isocaproic acid Mono-carboxylic CH₃ CHCH₂CHNH₂COOH mono-amino acids. Isoleucine or α-amino-β-methyl β-ethyl propionic acid CH₃ СНСНИН₂СООН Serine or a-amino B-hydroxy propionic acid CH_OHCHNH,COOH Aspartic * or a-amino-succinic acid COOH CH2CHNH2COOH Dicarboxylic Glutamic * or α-amino-glutaric acid mono-amino acids. COOH CH, CH, CHNH, COOH

*The amides corresponding to these two acids, namely asparagine CONH₂CH₂CHNH₂COOH and glutamine CONH₂CH₂CH₂CHNH₂COOH are of considerable importance in plants. The former occurs in asparagus and is produced in seeds which are allowed to germinate in the dark (Schulze, "Landwirtsch. Gahrb.," 1878, 411), while the latter has been found in the seeds of Cucurbita and many other plants (Schulze and Barbieri, "Ber. deut. chem. Gesells.," 1877, 10, 199; Schulze, id., 1896, 20, 1882). Asparagine and glutamine being readily hydrolysed by mineral acids, are not obtained as cleavage products of proteins by the ordinary methods of chemical hydrolysis, and for this reason are not quoted in the above list of cleavage products.

Ornithine or a-8-di-amino-valeric acid NH₂CH₂CH₂CH₂CHNH₃COOH Arginine or δ-guanidine α-amino-valeric acid Mono-carboxylic HN = Cdi-amino acids. NH CH, CH, CH, CHNH, COOH Lysine or α-ε-di-amino-caproic acid NH2CH2CH2CH2CH2CHNH2COOH Di-amino-trihydroxy-dodecanic acid * C10H26O5N2 Cystine or di [\$\beta\$-thio a-amino propionic] acid CH2-S-S-CH2 Dicarboxylic di-amino acid.

(2) Aromatic Compounds.

Mono-carboxylic mono-amino acids. Phenyl alanine or β-phenyl α-amino propionic acid C6H5CH5CHNH5COOH Tyrosine or β-parahydroxyphenyl α-amino-propionic acid HOC, H, CH, CHNH, COOH

(3) Heterocyclic Compounds.

Proline or α-pyrrolidine carboxylic acid CH.

Hydroxyproline or hydroxy α-pyrrolidine carboxylic acid Histidine or β-imidazol α-amino propionic acid

Tryptophane β-indole α-amino propionic acid C—CH₂CHNH₂COOH C₆H₄ CH

The above list comprises most of the more important cleavage products of proteins, the constitution of which has been definitely established.

Since different proteins give rise to different amounts of these various substances, it is obvious that a careful quantitative determination of the amounts of these acids produced by the hydrolysis of different proteins must be of considerable value.

^{*} The constitutional formula of this substance has not yet been determined

To this end Fischer, in 1901, introduced his so-called "Ester method," which consisted in converting the mixed amino acids obtained by hydrolysis of proteins into their corresponding esters, and then separating these by fractional distillation.

The method* is best illustrated by an example. Casein was decomposed by hydrolysis with concentrated hydrochloric acid, the hydrochloride of glutamic acid being separated by The filtrate was then evaporated under reduced pressure, taken up with alcohol and saturated with dry gaseous hydrogen chloride; in order to remove the water formed by the reaction, the solution was once more evaporated down, and the residue taken up with alcohol and again saturated with hydrogen chloride. The esters were next liberated from their hydrochlorides by evaporating the solution down to a syrup in a vacuum, diluting with water and approximately neutralizing by means of strong caustic soda solution while keeping thoroughly cooled in a freezing mixture. Concentrated potassium carbonate was now added, and the esters of aspartic and glutamic acid were extracted by ether; after adding more 33 per cent caustic soda and potassium carbonate and extracting again with ether, the combined extracts were dried with anhydrous sodium sulphate, evaporated and distilled under 8-15 mm. pressure. The various fractions were then separately hydrolysed, either by boiling with water or by warming them on the water bath with 20 per cent baryta water

This method, with slight modifications, has been applied by several workers, more especially Abderhalden and Osborne, to a considerable number of different proteins, with the result that there are now more or less reliable data for comparing the composition of proteins from various sources, both animal and vegetable.

A second method for gaining some insight into the composition of proteins consists in studying the distribution of nitrogen in the molecule with a view to ascertaining whether it is present in the form of mono- or di-amino acids, etc. A method for distinguishing between the different types of nitrogen-linking occurring in the molecule was first suggested by

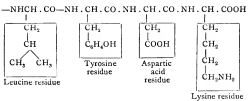
^{*} Fischer: "Zeit. physiol. Chem.," 1901, 33, 151.

Hausmann,* and has since been modified by Gümbel;† it depends on the fact that di-amino acids, in virtue of their strongly basic character, are precipitated from solution by the addition of phosphotungstic acid, whereas mono-amino acids are not.

The substance to be examined is first hydrolysed by boiling with concentrated hydrochloric acid for several hours under a reflux condenser. The amount of amide nitrogen and ammonia in the resulting mixture is then determined by distillation with magnesia in vacuo at 40° C.

- 2. The di-amino acid nitrogen is next determined by precipitating the residue in the flask with excess ‡ of phosphotungstic acid and estimating the amount of nitrogen in the precipitate by Kjeldahl's method (see p. 340).
- 3. The nitrogen combined as mono-amino acids may be determined directly in the filtrate or by the difference between the total nitrogen and the sum of the nitrogens separately determined by the above methods.

The fact that proteins on hydrolysis yield such a large number of amino acids, all of which have the amino group attached to the α -carbon atom (i.e., the carbon atom adjacent to the carboxyl), has led to the conclusion that the protein molecule is really composed of a long chain of these acids linked together in some such way as is represented below.



Such a compound would, of course, give the biuret reaction and contain but few free carboxyl groups or amino groups, which is entirely in agreement with the properties of proteins. Acting on this assumption, Fischer has synthesized a number of compounds containing such a structure, with the object of studying their properties and comparing them, if possible,

^{*} Hausmann: "Zeit. physiol. Chem.," 1899, 27, 95; 1900, 29, 136.

⁺ Gümbel: "Beitr. chem. Phys. u. Path.," 1904, 5, 297.

[#] In order to ensure complete precipitation of arginine.

with natural proteins. To these synthetic substances he has given the general name of Polypeptides.

The simplest polypeptide known is glycylglycine; this substance was obtained as follows:—

Glycine, when kept for some time in aqueous solution, loses water from two molecules, giving an anhydride

This substance, when boiled with hydrochloric acid, is hydrolysed, the ring being opened with the formation of the dipeptide glycylglycine.

$$\begin{array}{c} \text{CH}_2\text{--CO} \\ \text{NH} \\ \text{NH} + \text{H}_2\text{O} = \text{NH} \\ \text{COCH}_2\text{NH}_2 \end{array}$$

To give anything like a complete account of the methods employed in the synthesis of polypeptides is outside the province of this book. It may, however, be mentioned that a very fruitful method of synthesizing these substances consists in acting on an amino acid or a polypeptide with chloracetyl chloride, thus:—

CH₂CICCCI₊NH₂CH₂CONHCH₂COOC₂H₅=CH₂CICONHCH₂CONICH₂COOC₂H₅+HCI The latter, after conversion into the acid, and treatment with ammonia, yields a tripeptide,

 $CH_2CICONHCH_2CONHCH_2COOH+NH_3=CH_2NH_2CONHCH_2CONHCH_2COOH+HC1-Diglycy[g]ycine~a~1ripeptide \\$

Another valuable method consists in treating an amino acid suspended in acetyl chloride with phosphorus pentachloride and so obtaining an acid chloride R₁CHNH₂COCl. This latter is then allowed to act upon the amino group of a second acid as follows:—

$$R_2$$
 R_3

 $R_1CHNH_2COCI + NH_2CH \cdot COOH = R_1CHNH_2CONH \cdot CHCOOH + HCI$

The resulting polypeptide may, of course, be of considerable complexity, according to the nature of $R_{\rm 1}$ and $R_{\rm 2}.$

By these and similar methods, employing other combina-

tions of amino acids, polypeptides containing a great many different groupings have been synthesized. The one with the longest chain as yet obtained is an octodecapeptide leucyltriglycyl-leucylcrtoglycyl-glycine of the formula—

 $\begin{array}{c} NH_2CHC_4H_9CO[NHCH_2CO]_3NHCHC_4H_9CO[NHCH_2CO]_3NHCHC_4H_9CO[NHCH_2CO]_8\\ NHCH-COOH \end{array}$

The more complex of these polypeptides resemble the proteins in being colloidal substances which give the biuret reaction, and in being precipitated from solution by phosphotungstic or tannic acids and by ammonium sulphate.

The action of digestive ferments upon them has been studied by Abderhalden and others; they are not readily attacked by pepsin, but are hydrolysed by pancreatic or intestinal juice.

A striking confirmation of Fischer's view concerning the close connexion existing between the polypeptides and the natural proteins is to be found in the fact that the hydrolysis of proteins, under suitable conditions, has yielded four substances which could be identified with synthetic polypeptides. Thus, a solution of silk fibroin in hydrochloric acid was allowed to stand for several days; on evaporating a residue was obtained which, when digested with trypsin, yielded a peptonelike substance; the latter on hydrolysis with barium hydroxide gave glycylalanine, which was identified by its naphthaline sulphonic acid derivative.* Subsequently, the hydrolysis was repeated under somewhat altered conditions, with the same result that glycylalanine was obtained.† In a later communication, the same authors described the isolation of glycyltyrosine from the products of hydrolysis of silk fibroin, and of glycyl-leucine from elastin. Levene and Beatty; also claim to have obtained prolyl-glycine from the hydrolysis of gelatine.

Abderhalden § also mentions certain substances of a polypeptide nature which he found amongst the products of pancreatic digestion of a number of proteins such as caseine, edestine, hæmoglobin, serum globulin, egg albumin and fibroin.

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* Fischer and Abderhalden: "Ber. deut. chem. Gesells.," 1906, 39, 752. † Fischer and Abderhalden: "Ber. deut. chem. Gesells.," 1906, 39, 2315. ‡ Levene and Beatty: "Ber. deut. chem. Gesells.," 1906, 39, 2060.
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[§] Abderhalden: "Zeit. physiol. Chem.," 1905, 44, 28, 33.

OCCURRENCE OF AMINO ACIDS IN PLANTS.

Leucine occurs as such in the buds of the horse-chestnut * and many other plants. Isoleucine has been discovered by Felix Ehrlich † in the residual molasses obtained from sugar refineries.

Lysine and histidine have been isolated from sprouting plants by Schulze.‡

Arginine has been observed in the cotyledons of lupin seeds and in etiolated pumpkin seeds, \$ and also in several species of conifers.

Phenyl alanine was discovered by Schulze and Barbieri in etiolated germinating lupin seeds.

Tyrosine, according to Shibata, \(\text{occurs} \) occurs in considerable quantity in rapidly growing shoots of Japanese bamboos, and in small quantity in seedlings of Lupinus albus** and Vicia sativa; †† it has also been described as occurring with asparagine in the root-tubers of Dahlia variabilis. According to Bertel, ‡‡ tyrosine is converted into homogentisinic acid by an oxidase (see p. 350) contained in the plant, which is a change similar to the one produced in the human body in the condition known as alkaptonuria. §§ Schulze and Castoro, || however, deny the accuracy of these observations.

Tryptophane has been found in seedlings of Lupinus albus, Vicia sativa, and in Pisum sativum, II

Proline is obtained by the hydrolysis of a number of proteins of vegetable origin, notably the prolamins, but has not so far been found to occur as such in any plants.

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*Schulze and Barbieri: "J. prakt. Chem.," 1882, 15, 145; Schulze and
Winterstein: "Z. physiol. Chem.," 1902, 35, 299.
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[†] Ehrlich: "Ber. deut. chem. Gesells.," 1904, 37, 1809.

[‡]Schulze: "Z. physiol. Chem.," 1899, **28**, 465. §Schulze and Steiger: "Z. physiol. Chem.," 1887, **11**, 43; "Ber. deut. chem. Gesells.," 1886, 19, 1177.

[[]Schulze and Barbieri: "Ber. deut. chem. Gesells.," 1881, 14, 1785.

[¶] Shibata: "J. Coll. Sci. Tokyo," 1900, 13, 329.

^{**} Schulze and Castoro: "Z. physiol. Chem.," 1906, 48, 387, 396.

[#]Gorup Besanez: "Ber. deut. chem. Gesells.," 1877, 10, 781.

[#] Bertel: "Ber. deut. bot. Gesells.," 19 -2, 20, 451-

^{§§} Wolkow and Baumann: "Z. physiol. Chem.," 1891, 15, 266.

^{|| ||} Schulze and Castoro: loc. cit.

^{¶¶} Schulze and Winterstein: "Z. physiol. Chem.," 1910, 65, 431.

CLASSIFICATION OF PROTEINS.

The classification of the proteins was originally, for want of chemical knowledge, based on their different physical properties, such as solubilities, coagulation by heat, precipitation by neutral salts, etc.

Now that, from a study of their products of hydrolysis, a little more is known of the chemistry of the proteins, it is found that, on the whole, the physical method of classification is more or less in accordance with the chemical evidence.

Protamines.—These are the simplest proteins known, and are represented by such substances as salmine, sturine, cyclopterine, etc., which have been isolated from fish sperm.*

They usually occur associated with nucleic acid in the form of salts.

No compounds resembling the protamines have as yet been isolated from plants, although they may possibly occur in pollen.

Histones.—The histones, of which the best known one is that obtained from blood corpuscles, are characterized by being precipitated from solution by ammonia; they are related to the protamines, but are more complex than these substances.

Albumins.—This group includes egg-albumin, serum-albumin, and such vegetable albumins as legumelin of the pea and leucosin of wheat and other cereals.

The albumins are typically *soluble* in water and are coagulated by heat. They are *not* precipitated by saturation with sodium chloride or magnesium sulphate, nor by *half* saturation with ammonium sulphate, but, like all proteins, are precipitated by complete saturation with ammonium sulphate.

Traces of albumins occur in practically all seeds, but no seeds, so far examined, have been found to contain large quantities.

While plant albumins resemble those of animal origin

^{*} Kossel: "Bull. soc. chim., Paris," 1903, [23], 29, I-XVIII.

in regard to the two essential features of this group, namely, solubility in water and coagulation by heat, they differ in regard to their behaviour towards strong solutions of inorganic salts. Thus animal albumins are not supposed to be precipitated by half saturation with ammonium sulphate or saturation with sodium chloride or magnesium sulphate, but this is not always found to be the case for vegetable proteins, many of which are precipitated under these conditions.

Globulins.—These are exemplified by serum globulin, fibrinogen, and myosinogen, and also the derivatives of the two latter, fibrin and myosin. Examples of vegetable globulins are furnished by conglutin from the seeds of Lupinus, edestin from the seeds of Cannabis sativa, excelsin from the seeds of Bertholletia excelsa, legumin from the seeds of Pisum sativum, Vicia Faba, and other Leguminosæ, juglansin from the seeds of Juglans spp., vicilin from the seeds of Pisum sativum, Vicia Faba, etc., and vignin from the seeds of Vigna sinensis. In brief, globulins are amongst the commonest protein reserves of the higher plants.

The typical globulins are *insoluble* in pure water and are coagulated by heat. They are soluble in dilute salt solutions, but are insoluble in stronger salt solutions; thus, unlike the albumins, they are precipitated by saturation with magnesium sulphate or by only half saturation with ammonium sulphate.*

The vegetable globulins, which form the major portion of the reserve proteins of all seeds except cereals, do not always conform to these conditions of solubility. Thus, whereas animal globulins are insoluble in water and are precipitated by half saturation with ammonium sulphate, a great many globulins from plants are precipitated at less than half saturation, and, on the other hand, some are not precipitated until the solution is almost saturated with

^{*} These differences in solubilities between albumins and globulins may be illustrated by dissolving some of the white of an egg in water and placing it in a dialyser; as the small quantity of sodium chloride contained in the egg-white diffuses out, the globulin is precipitated out of solution; or again, if the solution is mixed with an equal volume of saturated ammonium sulphate solution, the globulin will likewise be precipitated out, owing to the solution now being half saturated with ammonium sulphate, but the albumin will remain in solution.

ammonium sulphate. It must, however, be noted that globulins extracted from seeds are nearly always obtained in the form of salts with a small amount of acid, and so long as they are in this form they have the characteristic solubilities of animal globulins. As soon, however, as the acid is removed they lose these and become completely soluble in water.

A further point of difference between animal and vegetable globulins is that many of the latter are only coagulated by heat with considerable difficulty.

The albumins and globulins are the only classes of proteins which are coagulated with heat.

Glutelins.—This is a small class represented by two proteins, both of vegetable origin, namely, glutenin found in wheat and oryzenin in rice. Similar substances probably occur in other cereals as well, but owing to the difficulty of obtaining them in a pure condition, they have not as yet been investigated.

Glutelins are insoluble in water and neutral saline solutions, but dissolve in dilute alkali or acid.

Gliadins or Prolamins.—These also are represented only by vegetable proteins, namely, gliadin from wheat or rye, hordein from barley, and zein from wheat or maize. Up to the present they have only been found to occur in cereals. The gliadins differ from all other proteins in being soluble in 70-90 per cent alcohol, the solutions being unaltered by boiling; they are insoluble in water or in salt solutions, but are soluble in dilute acids or alkalis.

On hydrolysis they yield a considerable quantity of proline (hence the name prolamins), glutamic acid and ammonia, but only small amounts of arginine and histidine, and no lysine.

Glutelins and gliadins are the chief protein constituents of the substance known as gluten.

Sclero-proteins.—This term is synonymous with the older term albuminoid, and includes substances of skeletal origin, such as keratin from hair, horn, etc., gelatin, elastin, and silk fibroin.

No representative of this class has as yet been found among vegetable proteins.

Phospho-proteins.—This group, which is probably not represented in the vegetable world, contains such substances as caseinogen and vitellin, obtained from milk and egg yolk respectively. The phosphorus of these proteins is in intimate organic combination with the protein molecule, and is not contained in the "prosthetic group" (see below) as in the case of the nucleo-proteins, which are composed of proteins with the phosphorus-containing nucleic acids.

The phospho-proteins are insoluble in water, but soluble in alkalis.

Caseinogen and vitellin were formerly known as nucleo-albumins, but the term is misleading, owing to the confusion arising with the nucleo-proteins, which are conjugated proteins (see below), and the term nucleo-albumin has for that reason been abolished.

The phospho-proteins resemble the nucleo-proteins in their solubilities, but they differ from them in their behaviour on hydrolysis; they yield at first a so-called pseudo- or para-nuclein, corresponding to the formation of a nuclein from a nucleo-protein, but whereas a nuclein on further hydrolysis yields nucleic acid, and ultimately purine bases, the pseudonuclein yields no corresponding pseudonucleic acid, but on the other hand is broken up by baryta water into phosphoric acid, but gives no purine bases.

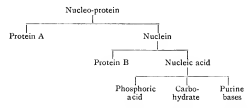
Conjugated Proteins.—This group may be divided into three sub-groups.

- I. Chromo-proteins, represented by hæmoglobin.
- 2. Nucleo-proteins, obtained from blood, chyle and lymph.
 - 3. Gluco-proteins, represented by mucin.

Conjugated proteins are characterized by the fact that on hydrolysis they break up, yielding a true protein and a substance of a different nature, for which Kossel has proposed the name "prosthetic" group.

Thus, for example, a chromo-protein like hæmoglobin breaks up into globin (a protein) and a pyrrole derivative, hæmatin (cf. chlorophyll, p. 236). Similarly, a glucoprotein such as mucin yields a protein and a carbohy-

drate, glucosamine.* And lastly, a nucleo-protein, when subjected to peptic digestion, or treated with dilute acid, gives a protein and a nuclein; this latter with caustic alkali breaks up still further into a second protein and a nucleic acid; the nucleic acids on further hydrolysis yield phosphoric acid, a carbohydrate residue, either a pentose or glucose, and purine bases, such as guanine, adenine, xanthine, etc. These changes are rendered clearer by the following scheme:—



It must be borne in mind that the protamines and histones frequently occur loosely combined with nucleic acids in the form of salts, but this type of combination is different from that between a protein and a nuclein such as is found in true nucleo-proteins.

The conjugated proteins appear to be rarely found in plants.

With regard to the occurrence of nucleo-proteins among plants, it is undoubtedly true that nucleic acid has been repeatedly found in plants, and compounds of proteins with nucleic acid have been isolated by Osborne, but it is not certain whether these substances actually occurred pre-formed in the seed, or were produced during the process of their isolation. Osborne† is

^{*} Glucosamine is a peculiar nitrogen containing sugar of the formula

CH₂OHCHOHCHOHCHOHCHNH₂CHO or CH₂OHCHOHCH CHOH CHNH₂CHOH It has all the ordinary reactions of sugars as regards reduction of Fehling's solution, reaction with phenylhydrazine, etc., but is not fermentable by yeast. Owing to the presence of the amino group, it is also able to form salts with acids such as hydrochloric acid. It was first obtained by the hydrolysis of chitin contained in the shell of lobsters and has since been obtained by the hydrolysis of several gluco-proteins such as serum mucoid, etc.

⁺ Osborne: "The Vegetable Proteins," London, 1909.

of opinion that "only small quantities of nucleo-protein occur in the entire seed, and that this will be found chiefly in the tissues of the embryo in which the nuclei of the cells are far more abundant than in the tissues of the endosperm".

With regard to chromo-proteins and gluco-proteins, the former possibly may be represented by phycoerythrin (p. 258) and the latter by the mucilage which occurs in the roots of *Dioscorea japonica*, which in many of its characters resembles mucin from animal sources.

Derivatives of proteins.—In this group are included a number of substances obtained by the hydrolysis of proteins; they may be sub-divided as follows:—

- I. Meta-proteins, consisting of acid albumin and alkali albumin, produced respectively by the action of acid or alkali on proteins.
- 2. Proteoses, represented by albumose, globulose, gelatose, etc. These substances are produced from proteins by the action of digestive juices such as pepsin and trypsin.

Pepsin, which acts in an acid medium, breaks up the protein as follows:—

PROTEIN.

META-PROTEIN (acid albumin).

PRIMARY PROTEOSE (precipitated by half-saturated ammonium sulphate and by potassium ferrocyanide in the presence of acetic acid).

Secondary Proteose (precipitated by saturated ammonium sulphate, but only slowly by potassium ferrocyanide in the presence of acetic acid) Peptone (not precipitated by saturated ammonium sulphate nor by potassium ferrocyanide in the presence of acetic acid). Polypeptides and Amno Acids.

POLYPEPTIDES AND AMINO ACIDS.

The formation of amino acids from peptones takes place only after prolonged action.

Trypsin, which acts in an alkaline medium, produces substantially the same series of changes, only that the meta-protein in this case is alkali albumin; furthermore the decomposition into amino acids takes place more rapidly than with pepsin.

A great many seeds have been found to contain proteoses after the removal of the other proteins, and substances resembling the proto- hetero- and deuteroproteoses obtained from animal proteins have been described; but in all cases it is difficult to say whether these substances were not produced by some secondary action of enzymes upon the protein, during the process of isolation

- 3. Peptones. Substances belonging to this class still give the biuret reaction, but unlike all other proteins they are not precipitated from solution by saturation with ammonium sulphate.
- 4. Polypeptides, which include such substances as leucyl glutamic acid, obtained by Fischer and Abderhalden from gliadin by hydrolysis with 70 per cent sulphuric acid, and glycyl tyrosine and glycyl leucine, obtained by the same authors from silk fibroin and elastin respectively.

COMPARISON BETWEEN VEGETABLE AND ANIMAL PROTEINS.

From the foregoing it will be seen that, in the main, the animal and vegetable proteins conform sufficiently well with regard to their general properties and solubilities that they may be included in the same scheme of classification. The greatest irregularities are exhibited in the groups of albumins and globulins, but even these are not sufficiently serious to suggest any fundamental difference between the proteins derived from animal and vegetable sources. These views are confirmed by chemical evidence: with the single exception of di-amino trihydroxy-dodecanic acid, a substance as yet only obtained from casein, all the known products of hydrolysis of animal proteins have been obtained from vegetable proteins, and there is no real reason for assuming that there is any fundamental difference in the structure of the protein molecule from the two sources.

On the whole, vegetable proteins yield more glutamic acid, and many also yield rather more proline, arginine and ammonia than do animal proteins.

The comparatively large quantities of proline and arginine which occur in some cases may be responsible for the slightly higher nitrogen content which characterizes proteins of vegetable origin.

Further, it should be noted that the prolamins, or alco-

hol soluble proteins, form a distinct class; they are found only in the vegetable kingdom, and have no analogues amongst proteins from animal sources.

Of twenty-three different seed-proteins which have so far been systematically hydrolysed, all were found to contain leucine, proline, phenylalanine, asparagine, glutamic acid, tyrosine, histidine, arginine and ammonia; two gave no glycine; two gave no alanine; four gave no lysine; and one gave no tryptophane. One, namely zaein, gave neither glycine, lysine nor tryptophane. Three gave no cystine, and two others only traces.

It is, on the whole, unlikely that there is any protein entirely free from sulphur, although in the case of vicilin the amount is actually as low as 0.1 per cent. If it is assumed that the sulphur is contained in the molecule in the form of cystine, it follows that there must be at least two atoms of sulphur present. Calculations based on this assumption give a value for the molecular weight of at least 15,000, but although from other considerations the molecular weight of proteins is known to be high, it is unlikely that the value is as high as this.

While it is possible by means of general reactions to place a given protein in the class of albumins or globulins, there are no distinctive chemical or physical methods by which the identity of any particular albumin or globulin may be established; thus it not infrequently occurs that two substances which have been obtained from different sources, and are described under different names, are eventually found to give the same figures on analysis, and are therefore regarded as identical. This is notably the case with albumins obtained from different plant seeds, and the serum albumin derived from different animals. Within the last few years, however, a biological method has been discovered which promises to become of the very greatest value in distinguishing the various compounds from each other. Following upon the researches of Wassermann and Uhlenhuth. Tstistowitch found that serum drawn from a rabbit, which had been inoculated for some time with the serum of a horse, had acquired the property of producing a precipitate when added to normal horse serum: this is due to the formation in the rabbit's blood of a substance known as a precipitin, which

belongs to a class of compounds described by Hofmeister as pseudo-globulins; the precipitate formed is a compound of the precipitin with the albumin contained in the serum to which the precipitin was added. The precipitin so prepared should only react with horse serum and not with the serum of any other animal; the reaction is, however, not absolutely specific, inasmuch as a precipitin may react with the serum of an animal closely related to the one from whose serum it was prepared. It has, moreover, been found that substances which had been isolated from natural fluids, as well as native sera, were able to incite a precipitin formation when injected into the blood of some living animal, and it has been thus possible to show that the albumin contained in milk is not identical with that obtained from blood. The method has been employed by Kowarski and Schütze* for distinguishing the various plant albumins, and by Rickmann, + Uhlenhuth, ± and others for distinguishing between horse flesh and the meat of other animals. An attempt has also been made to employ the same principle for the estimation of proteins by a comparison of the precipitates formed under various conditions.§

As illustrating the very close connexion existing between albumins and globulins, it is worthy of note that Moll claims to have converted serum albumin into serum globulin by warming a 3 per cent solution of serum albumin for one hour to 60° C. with N/66 sodium carbonate, but it is difficult to say whether true serum globulin was actually produced. According to Chick and Martin, || however, the conversion of albumin into globulin may be explained merely by assuming a difference in the state of aggregation.

EXTRACTION OF PROTEINS.

The main facts relating to the solubilities of the common vegetable proteins are as follows:-

^{*} Kowarski and Schütze: "Deut. med. Wochenschr.," 1901, 27, 442; 1902, 28, 804.

[†] Rickmann: "Chem. Zeit.," 1907, 2, 1983. ‡ Uhlenhuth and Weidanz: "Praktische Anleitung z. Ausführung des biolog. Eiweissdifferenzierungsverfahren," Jena, 1909.

[§] Schulz: "Deut. med. Wochenschr.," 1906, 32, no. 26; "Zeit. Unters. Nahr. u. Genussm.," 1906, 12, 257.

| Chick and Martin: "Journ. Physiol.," 1912, 45, 261.

- 1. Proteoses, albumins, and some globulins are soluble in water.
- 2. Globulins, together with most of the proteins soluble in water, dissolve in 10 per cent sodium chloride.
 - 3. Prolamins are soluble in alcohol (70 to 90 per cent).
- 4. Glutelins and prolamins dissolve in dilute acid and in dilute alkali.

These facts are made use of in the extraction of the substances in question from vegetable tissues such as seeds, which may contain several proteins; and although the products so obtained are anything but pure, a brief outline of the method may be given. The separation of the proteins removed by these means from the seed by a given solvent is a very lengthy and tiresome process, and the details must be sought for elsewhere.*

Before proceeding with the extraction, the material must be ground up as finely as possible, in order that all the cells may be broken; if needs be, the tissue must be carefully dried beforehand, but too high a temperature must not be used.

In all cases the initial procedure is much the same; the main point to be observed, as in everything else, is thoroughness. The powdered material is well mixed with the solvent, which is allowed to act for some time; the mixture should be well shaken periodically. The solid is then filtered off and well washed with fresh solvent, and is again treated until the extract gives no protein reaction. The temperature may be raised during the extraction, but it should not be high enough to alter the proteins. If the extraction, especially with aqueous solvents, be prolonged, it may be necessary to add a little antiseptic, such as chloroform, in order to stop bacterial action.

When it is desired to make successive extracts, in cases such as seeds where several proteins may be present, the order may be water, 10 per cent sodium chloride, alkali ('1 to '2 per cent caustic potash or '5 to 1 per cent sodium carbonate), and finally alcohol (70 to 80 per cent).

The initial extraction may be made with salt solution, the albumins being afterwards separated from the globulins, and

^{*} See Osborne: "The Vegetable Proteins," London, 1909, on whose account the following is based. For a method for the preparation of plant globulins, see Reeves: "Biochem. Journ.," 1915, 9, 508.

this course is recommended when both are present on account of the saving of time.

The proteins isolated by these means may be roughly purified as follows:—

- 1. Albumins and globulins.—These will nearly always be contaminated one with the other. A separation may be effected in the following ways:—
 - (a) The solution is saturated with magnesium sulphate, whereby the globulins are precipitated and the albumins remain in solution (but see above, under albumins and globulins).
 - (b) Dialysis. The extract is placed in a dialyser and floated in water which is continually changed. The precipitated globulins are filtered off from the salt solution, which, of course, is getting weaker and weaker and contains the albumins. The precipitated globulins are re-dissolved in warm saline solution, which may on cooling deposit globulins in a crystalline form; if this does not occur, the solution may be saturated with magnesium sulphate. The albumins may be precipitated by saturating the solution with ammonium sulphate. Further purification may be effected by a repetition of the process and by fractional precipitation with magnesium sulphate or by ammonium sulphate, according to the protein to be purified.
- 2 Glutelins.—The proteins soluble in dilute alkali may be precipitated by very carefully neutralizing the solution and then further adding only just sufficient acid to cause the precipitation of the glutelins. The precipitate may be well washed with a dilute neutral saline solution, in which it is insoluble, in order to remove any globulins which may be present.
- 3. Prolamins.—The extract, which is made by treatment with hot alcohol, is either mixed with water sufficient in amount to precipitate the proteins, or the filtered solution may be evaporated under a reduced pressure at a temperature not higher than 50° C. The precipitate is filtered off and may be re-dissolved in as little alcohol as possible. From this solution the protein may be recovered by the addition of absolute

alcohol, in which prolamins are insoluble, and ether. The ether is added in order to make the precipitation more complete and also to hold any fats which may have been extracted by the alcohol.

THE SYNTHESIS OF PROTEINS IN THE PLANT.

In view of our limitations as regards the chemistry of proteins, it is not surprising that we are in almost complete ignorance respecting the synthesis of these substances in plants.

It is generally agreed that the leaves are the important centres of protein formation, and they show a periodicity in their nitrogen content. Thus Otto and Kooper * and Le Clerc du Sablon—found that there is a gradually decreasing amount of nitrogen from the spring to the autumn, and that leaves of several different plants, even in different stages of development, exhibit a greater nitrogen content in the morning than in the evening.

The requisite nitrogen is obtained, not from the air—the Leguminosæ are here excluded—but from the salts contained in the water absorbed by the roots; thus the fertility of soil, especially with regard to nitrates, is most important, as has been shown by direct experiments.‡

In passing, it may be remarked that some plants, at any rate, can make use of salts of ammonium as a source of nitrogen. Hutchinson and Miller § found this to be true under conditions of culture which precluded the presence of nitrates in the soil. In this respect, however, all plants do not behave alike; whilst some will grow equally well whether supplied with nitrates or ammonium salts, others flourish best when supplied with the former, and others seemingly prefer ammonium salts to begin with and then nitrates.

Since protein formation takes place particularly in the leaves, it might be supposed that light is an important and direct factor in its synthesis. Indeed earlier workers, Schimper || for example, considered this to be so, but more recent investigations tend to show that the synthesis of proteins

^{*}Otto and Kooper: "Landwirthsch. Jahrb.," 1910, 39. + Le Clerc du Sablon: "Rev. Gen. Bot.," 1904, 16, 341. ‡Whitson and Stoddart: "Ann. Rep. Wisconsin Exp. Sta.," 1904, 193.

[§] Hutchinson and Miller: "Journ. Agric. Sci.," 1909, 3, 179.

[|] Schimper: "Flora," 1890, 73, 207.

can take place in the dark and in tissues free from chlorophyll, provided that an adequate supply of carbohydrate be at hand.* Zaleski and Suzuki† found that the leaves of the sunflower floating upon a solution containing sugar and nitrate produced considerable quantities of proteins in the dark, from which it appears that nitrate assimilation is not a photochemical process, and that light is only of indirect importance in providing one of the means for the formation of carbohydrates.‡ The synthesis of proteins is conditioned by the available supply of carbohydrate, and since photosynthesis is a daylight process, it is not surprising to find that the production of proteins may be four or five times as great in the light as in darkness. \ Baudisch || is of the opinion that the formation of protein under abnormal conditions in the dark is no proof that the process is not a photochemical one under normal conditions: he considers that the synthesis may in this case be due to anærobic respiration or some other abnormal chemical processes which reduce the nitrates and so aid in the production of proteins. It has also been stated that if but small quantities of carbohydrate are available, the synthesis of proteins, in darkness, may stop at the formation of amides, which some plants, e.g., Algæ such as Pleurococcus and Raphidium, and the Fungi Eurotium and Penicillium, can directly assimilate.**

Evidence is not wanting to show that nitrates are reduced to nitrites in the plant,†† but much uncertainty exists as to the further fate of the nitrite. To obtain some insight into this

^{*} Jost: "Biol. Centrlbl.," 1900, 20, 625.

[†]Zaleski and Suzuki: "Ber. deut. bot. Gesells.," 1897, 15, 536; "Bot. Centrlbl.," 1901, 87, 281; Suzuki, "Bull. Coll. Ag. Tokyo," 1898, 2, 409; 3, 241.

[‡]Zaleski: "Ber. deut. bot. Gesells.," 1909, 27, 56.

[§] Montemartini; "Atti. R. Inst. Bot. Pavia," 1905, II, 10, 20.

[|] Baudisch: "Zentr. Bakt. Parasit.," 1912, 32, 520. | ¶ Jakobi: "Biol. Centrlbl.," 1898, 18, 593.

^{**} Lutz: "Bull. Soc. Bot. France," 1902, 48, 118.

^{††}Pertiabosco and v. Rosso: "Staz. sperim. Agrar. Ital.," 42, 5; Aso: "Beih. bot. Centribl.," 1900, 15, 208.

Since plants absorb their nitrogen chiefly in the form of nitrates, there should be some means of reducing them to nitrites. According to Irving and Hankinson ("Biochem. Journ.," 1908, 3, 87) such an enzyme occurs in many plants, chiefly aquatic, examined by them. These plants gave off nitrogen; but oxygen, owing to the reducing agent, was not evolved. The sequence of events which they suggest is as follows:—

question Baudisch * conducted experiments outside the plant, to ascertain the effect of light upon nitrates and nitrites. He found that potassium nitrate, on exposure to diffuse daylight, loses oxygen, and likewise potassium nitrite mixed with formaldehyde or methyl alcohol was, by exposure to diffused daylight, reduced to potassium hyponitrite; the latter compound, however, combined with formaldehyde to give the potassium salt of form-hydroxamic acid—

$$CH_3OH + KNO_2 = OHCH : NOK + H_2O$$

Prolonged exposure to light caused the reduction to ammonia, and led to the production of potassium carbonate; the following gases were also evolved, carbon dioxide, carbon monoxide, oxygen, nitrous oxide and hydrogen,—an element which has been stated to be given off by plants.† As a result of these experiments, Baudisch comes to the conclusion that in the plant ammonia is oxidized by oxidases or by ultra-violet light; the resulting product then combines at once with formaldehyde to give aci-nitromethane $\mathrm{CH_2} = \mathrm{NOOH}$, a substance isomeric with formhydroxamic acid. Aci-nitromethane is known to be very reactive, and it is shown by a number of equations to be capable of giving rise to several more or less complex substances frequently found in plants.

The hydrolysis of proteins yields amino acids, such as leucine, asparagine, and tyrosine, which also occur in protein-containing seeds. If the stages in the destruction and construction of proteins correspond, i.e. if the action be reversible, then the synthesis of amino acids becomes the first problem; the second is the linking together of these substances to form the finished product. Zaleski‡ considers that the action is reversible and that the amino acids formed in the hydrolysis of proteins are also involved in their synthesis. He found

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 \begin{array}{cccc} \text{I. } 2\text{KNO}_3 \rightarrow 2\text{KNO}_2 + \text{O}_2 \\ \text{2. } \text{KNO}_2 \rightarrow \text{HNO}_2 \text{ (by the activity of the acid of the cell sap)} \\ & \text{CH·NH}_2\text{-COOH} & \text{CHOH·COOH} \\ \text{3. } 2\text{HNO}_2 + \Big|_{\text{CH}_2\text{-CONH}_2} \rightarrow \Big|_{\text{CH}_2\text{-COOH}} + 2\text{H}_2\text{O} + 2\text{N}_2 \\ & \text{Asparagine} \end{array}
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^{*} Baudisch: "Ber. deut. chem. Gesells.," 1911, 44, 1009.

[†]Polacci: "Atti. Inst. Bot. Pavia," 1902, 7, 97.

‡Zaleski: "Ber. deut. bot. Gesells.," 1905, 23, 126; "Beih. bot. Centralbl.," 1911, 27, 63.

that during the ripening of pea seeds there was an increase in the amount of the protein at the expense of the amino acids and organic bases, as indicated by nitrogen determinations of these compounds.

	Control.			After five days.			
N of proteins N of amino acids . N of organic bases . N of other compounds		79°2 p 8°7 10°8 1°4	er cent	of total N	89.2 p 4.6 5.6	er cent	of total N

This synthesis is hastened by drying and the presence of free oxygen, which gas, however, has an indirect action.

The results were not so well marked for all seeds; thus under similar conditions but little protein synthesis took place in the maize, whilst in the sunflower there was a decomposition of protein.

Since the hydrolysis of proteins is naturally effected by means of enzymes, the question arises, is the action of these catalysts reversible? (See p. 363.)

The problem is the more complicated since substances may be formed in plants which do not occur when proteins are hydrolysed in the test tube. Asparagine is a case in point. This substance is sometimes very abundant in plants, especially in seedlings germinated in the dark,* and if it be a direct stage in the analysis of proteins, then it would appear that the sequence obtaining in the plant is not the same as that which happens in vitro. This, of course, is not impossible—indeed, some would say that it is extremely probable—but the opinion is now generally held that asparagine is a secondary product.† Thus it occurs in greater abundance in the developing parts than in the organs where the protein reserves are stored; Schulze‡ found that only 7.62 per cent of asparagine occurred in the cotyledons, whilst 31.81 per cent obtained in the axis of the lupin. Also the relative

^{*}This may easily be shown by germinating lupin seeds in the dark until the hypocotyl is a few inches in length. On mounting a section of the hypocotyl in strong alcohol and examining under the microscope, a large number of crystals of asparagine will be seen.

[†] Prianischnikow: "Ber. deut. bot. Gesells.," 1904, 22, 35; Schulze: id., 1904, 22, 385; 1907, 25, 213; Zaleski: id., 1906, 24, 292.
† Schulze: "Landw. Jahrb.," 1878, 411.

amounts of asparagine and aspartic acid show considerable variation during germination and, in the last stages, the amount of asparagine formed is in a proportion greater than the amount of protein decomposed.

The significance of asparagine to the plant is not known; it would not appear to be used up directly for the synthesis of new protein since its amount does not decrease proportionately to the quantity of protein which is built up; it has been suggested that asparagine is merely an intermediate step in the synthesis of carbohydrates or fats.

The method of formation of asparagine is uncertain; ammonia, which is formed during the decomposition of the primary dissociation products of proteins, may be, according to Schulze,* its starting point, and this combines with an amino acid, possibly aspartic acid, according to Prianischkow, to form ammonium aspartate, which by the loss of a molecule of water gives origin to the asparagine.

Other views regarding protein synthesis have been put forward;† but owing to their not being based on well ascertained facts, it does not seem profitable to discuss them here.

Treub‡ concluded, from his investigations on the distribution, periodic variation in the amount, etc., of cyanogenetic glucosides (see section on glucosides), that hydrocyanic acid is the first recognizable product of nitrogen assimilation and possibly is the first organic nitrogen compound formed, but the conclusions are not convincing. On purely chemical grounds it is not impossible that acetonecyanhydrin CH₃COHCNCH₃ may be a stage in protein synthesis.

From the foregoing it is obvious that our knowledge of the synthesis of proteins in the plant is in a very unsatisfactory condition; and this need cause no surprise in view of the enormous difficulty of the subject and the extraordinary complexity of the protein molecule.

SYNTHESIS OF AMINO ACIDS IN THE PLANT.

With regard to the synthesis of amino acids within the plant, it is of interest to note that in the laboratory Erlenmeyer

^{*} Loc. cit. See also Treboux: "Ber. deut. bot. Gesells.," 1904, 22, 570.

[†] Loew: "The Energy of Living Protoplasm," London, 1896, and Munich, 1899.

[†]Treub: "Ann. jard. bot. Buitenzorg," 1895, 13, 1, and 1904, 19, 86.

and Kunlin* have been able to synthesize the acetyl and formyl derivatives respectively of alanine and glycine by the action of ammonia on glyoxylic acid, both of which substances are known to occur in plants. The changes involved may be represented by the following formulæ:—

Furthermore, Fischer \dagger has been able to synthesize a diamino acid by the action of ammonia on sorbic acid, an unsaturated acid occurring in the unripe berries of the mountain ash; also another unsaturated acid belonging to the same series as sorbic acid, namely, β -vinyl acrylic acid, has by the action of ammonia been converted into diamino valeric acid, and further, aspartic acid \S has been obtained by the action of ammonia on fumaric acid.

From the plant physiological point of view, however, the interest of these latter discoveries is dependent on the occurrence in the plant both of unsaturated acids and of ammonia.

The researches of Ehrlich || upon the action of yeast on amino acids have led to some very interesting results; it was found that the addition of leucine or isoleucine to a fermenting sugar solution gave rise to a production of inactive or active amyl alcohol respectively, according to the following schemes.

$$\begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{CH}_{3} \\ \text{CH}_{2} \\ \text{CH}_{2} \\ \text{CHNH}_{2} \\ \text{COOH} \\ \text{Hu}_{2} \\ \text{COOH} \\ \text{CH}_{3} \\ \text{CHCH}_{2} \\ \text{CH}_{2} \\ \text{CH}_{2} \\ \text{CH}_{2} \\ \text{CH}_{2} \\ \text{CH}_{2} \\ \text{CH}_{2} \\ \text{CH}_{3} \\ \text{CHCH}_{2} \\ \text{CH}_{4} \\ \text{OH} \\ \text{COOH} $

The amounts of these alcohols produced are proportional

^{*} Erlenmeyer and Kunlin: "Ber. deut. chem. Gesells.," 1902, 35, 2438.

⁺ Fischer and Schlotterbeck: id., 1904, 37, 2357.

[‡] Fischer and Raske: id., 1905, 38, 3607.

[§] Engel: "Compt. rend.," 1887, 104, 1805, and 1885, 106, 1677.

^{||} Ehrlich: "Ueber die Bedeutung des Eiweissstoffwechsels, etc. " "Sammlung chem. u. chem. tech. Vorträge," Stuttgart, 1911.

to the quantities of leucine or isoleucine added and rise, under favourable conditions, to as much as 7 per cent; furthermore, it was found that although the leucine parted with its nitrogen in the form of ammonia, the latter substance was not lost, but appeared to be taken up by the yeast in the production of new protein material; this observation led to trying the effect of adding ammonium salts, when it was found that the yeast, finding these latter to be an easier source of nitrogenous food, gave up attacking the leucine, and consequently less amyl alcohol was produced.

These experiments therefore prove that amino acids can be fermented by yeast with the production of alcohols in much the same way as sugars can be fermented. The practical importance of these discoveries can be gauged from the fact that the production of amyl alcohol* or fusel oil by the yeast fermentation of sugar has always been a source of trouble to spirit distillers, and necessitated elaborate processes for refining; these researches have provided both an explanation of the cause and a remedy for the evil.+

Since, moreover, other amino acids besides the leucines are also found to be attacked in a similar way with the production of a number of widely different products, some of which are aromatic, it is easy to account for the different flavours which are peculiar to the various alcoholic beverages, all of which are ultimately produced by alcoholic fermentation of sugars in presence of different proteins.

The destruction of amino acids by enzymes derived from yeasts, fungi or bacteria with the formation of different byeproducts, may also account for the flavours of different cheeses, as well as the odour of flowers; the substance phenyl ethyl alcohol, for example, which was produced by the fermentation of phenyl alanine,

 $C_6H_6CH_2CHNH_9COOH + H_9O = C_6H_5CH_9CH_9OH + CO_9 + NH_3$ Phenyl alanine Phenyl ethyl alcohol

being the chief odoriferous constituent of rose oil.

* Besides these alcohols, other substances, such as succinic acid, glycerin, etc., are produced, but they also probably owe their origin to amino acids.

† A knowledge of the cause of the amyl alcohol production is also important from the point of view of increasing the yield of this substance, since large quantities of amyl alcohol are required for the preparation of amyl acetate, used as a flavouring material for confectionery, and as a solvent in the manufacture of varnish, smokeless powder, etc.

These researches would therefore lead to the conclusion that the proteins, through the breaking up of various amino acids derived from them, are ultimately responsible for the production of a variety of nitrogen-free alcohols, aldehydes and acids as bye-products, which go to produce the different essential oils, etc.

The metabolism of proteins in the animal world is, as is well known, a very important process and results in their very complete decomposition with the formation of urea, carbon dioxide and water. Although little is known concerning the metabolism of proteins by plants, there is good reason for believing that the destruction of the protein molecule is far less complete; the occurrence of urea has in fact only rarely been recorded in plants, namely, in Lycoperdon Bovista* and traces have been isolated from Cichorium endiva, Cucurbita maxima, Cucumis melo, Brassica olereacea and B. napus, Solanum tuberosum and Spinacia olereacea.† It has been suggested that many of the simpler nitrogenous compounds, as, for example, caffeine, theobromine, the alkaloids, skatol and allied substances such as indigo, indoxyl, etc., may be products of protein metabolism

ESTIMATION OF NITROGEN.

The Kjeldahl process for the estimation of nitrogen depends on the fact that organic nitrogenous compounds when boiled with concentrated sulphuric acid are decomposed with the formation of ammonium sulphate,‡ while the carbon and hydrogen are oxidized to carbon dioxide and water respectively by the sulphuric acid. In order to ensure complete destruction of the organic matter, the heating must be continued until the mixture is colourless, or, at most, light straw-coloured. This

† Fosse: "Compt. rend.," 1912, 155, 851; 1913, 156, 567, 1938; 1914, 158, 1374; 159, 253; Verschaffelt: "Pharmaceut. Weekblad," 1914, 51, 189.

^{*} Bamberger and Landsiedl: "Monatshefte," 1903, 24, 218.

[‡]Organic compounds containing the nitro- or nitroso-groups and also inorganic nitrates and nitrites, if heated with concentrated sulphuric acid, would give off oxides of nitrogen which would escape without conversion into ammonium sulphate; this difficulty may, however, be overcome by adding to the substance from 1-2 grams of zinc dust, and then rapidly pouring over the mixture, so as to cover it at once, a solution of 2 grams of salicylic acid in 20-30 c.c. of concentrated sulphuric acid, carefully heating until frothing ceases, and then proceeding with the addition of potassium sulphate, etc. The nascent hydrogen produced by the zinc and the acid probably reduces the nitro group to the amino group, which is then readily converted into ammonium sulphate.

may in some cases require prolonged heating, but the decomposition is usually accelerated by (a) adding potassium sulphate to the sulphuric acid in order to raise its boiling point, or (b) adding a catalytic agent in the form of a globule of mercury or some copper sulphate, or (c) adding an oxidizing agent, such as potassium permanganate.

When the decomposition is complete, the acid is cooled and diluted largely with water and finally made alkaline and distilled, the liberated ammonia being absorbed in a known quantity of standard acid; by determining the amount of free acid remaining, the amount neutralized by the ammonia is calculated, and hence the amount of nitrogen contained in the original substance may be deduced.

The actual experiment is carried out as follows:-

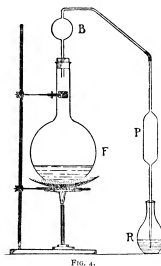
From 0.3 to 0.7 gram * of substance is weighed out into a 250 c.c. hard glass round-bottomed flask; 10 grams of pure potassium sulphate and a small globule of mercury, or a small quantity of mercuric sulphate, are added, and the mixture covered with 20 c.c. of pure concentrated sulphuric acid free from nitrogen; a small funnel is placed in the neck of the flask, which is then clamped in an inclined position and heated over an iron wire gauze in a fume cupboard. When the liquid is only faintly coloured, it is cooled and diluted with 150 c.c. of water and transferred to a 750 c.c. flask (Fig. 4); the small flask is then thoroughly rinsed, in small portions at a time, with another 150 c.c. of water, all of which are carefully poured into the large flask without loss.

Fifty c.c. of a standard acid, approximately decinormal, are now delivered into the flask R, a couple of drops of methyl orange are added, and the flask is stood in a vessel of cold water.

About 10 c.c. of a 10 per cent solution of potassium sulphide together with a few lumps of granulated zinc† and two or three pieces of litmus paper are next added to the diluted solution in the flask F. Sufficient 50 per cent caustic soda (usually about 50 c.c.) is now added to render the thoroughly mixed contents alkaline to the litmus paper; the flask is then

^{*} Sufficient substance should be taken to contain about 0.05 gram of nitrogen. †This when heated in the presence of alkali gives off a steady stream of hydrogen, and so prevents bumping.

connected without delay to the bulb tube B, to which is



attached the pipette P, whose lower end is then dipped into the acid in R.

The flask is then boiled over a sand bath until about 100 c.c. of liquid have distilled over.* Great care must be taken not to allow the flask F to become cool by exposure to draughts or by temporarily lowering or removing the flame, as otherwise the acid in R will rush back and spoil the experiment.

When the distillation is complete the cork in the large flask F is taken out and the flame is then removed; the pipette P next is disconnected from

the bulb tube B and carefully washed with water before withdrawal from the flask R. The excess of acid is then titrated back with standard alkali until the pink colour of the solution changes to orange.

The method of calculating the result may be seen from the following example:—

Weight of substance taken = 0.55 gram. Volume of .1045 N acid taken = 50 c.c.

Volume of '0946 N alkali required for back titration = 15.4 c.c.

50 c.c. \cdot 1045 N acid = 50 \times \cdot \cd

Volume of acid neutralized 3.769 c.c. N acid. But 3.769 c.c. N acid are equivalent to 3.769 c.c. N nitrogen. $\equiv \frac{3.70 \circ \times 14}{1000} = .0427 \text{ gram nitrogen}$

Per cent nitrogen =
$$\frac{100 \times .0427}{.55} = 7.76$$

* It is important to observe that the distillate should remain acid throughout the distillation as indicated by its pink colour; if at any time it should change to yellow, another 50 c.c. of standard acid should be added at once, and this must of course be taken into consideration in working out the result.

The amount of protein corresponding to the percentage of total nitrogen estimated in this way may be found by multiplying the percentage by the factor 6'37.*

*Assuming the percentage of nitrogen in protein to be, on an average, 15.7, the amount of protein corresponding to a given weight of nitrogen x will be

$$x \times \frac{100}{15.7} = x \times 6.32$$

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SECTION IX.

ENZYMES.

It has long been known to chemists that the velocity of chemical reactions could, in many cases, be increased by the presence of relatively small quantities of certain substances which do not appear to take any immediate part in the reaction.

This is well illustrated by the familiar example of the effect of a small quantity of manganese dioxide in bringing about the liberation of oxygen from potassium chlorate at a temperature much lower than would be possible by heating this substance alone.

Other examples of the accelerating influence of foreign substances on the velocity of reactions are to be found in the use of cuprous chloride in Deacon's chlorine process, and of spongy platinum, either in the manufacture of sulphuric acid by the contact process, or for effecting the explosive combination of hydrogen and oxygen.

Similarly, the hydrolysis of cane sugar according to the equation—

$$C_{12}H_{22}O_{11}\,+\,H_2O\,=\,2C_6H_{12}O_6$$

takes place very slowly in neutral aqueous solutions, but may be greatly accelerated by warming the solution with a little mineral acid.

A feature common to all the above reactions is the fact that the substance which produces the accelerating influence is unaltered by the reaction, and can usually be recovered from the reaction-product unchanged in quality and quantity.

Substances which have this remarkable property of being able in some way to influence the velocity of a reaction, without apparently undergoing any change themselves, and which act in quantities which bear no particular relation to the weights of the reacting substances, are called catalytic agents.

The process of catalysis has been defined by Ostwald as "The acceleration of a chemical change by the presence of some foreign substance," and it must be clearly understood that a catalytic agent only accelerates a reaction, but is not capable of bringing about a reaction which would not take place at all in its absence. Berzelius,* in 1850, drew attention to the similarity between the decomposition of hydrogen peroxide, under the influence of insoluble inorganic catalysts such as platinum or silver, and the decomposition of sugar into alcohol and carbon dioxide under the influence of substances known as ferments. Thus, in view of the case with which so many complex reactions are effected within the living organism at a low, or a comparatively low, temperature, the idea is suggested that nature likewise makes use of catalysts.

As a matter of fact a large number of complex organic substances, capable of exerting catalytic action, have been isolated from plants and animals; and to these substances the name of enzymes has been applied.

The food of plants, carbohydrate, protein, fats, etc., is, in many cases, valueless unless it can be brought into a condition suitable for assimilation and, very often, translocation. Thus the starch in a leaf must be rendered soluble before it can be transported to other parts of the plants, and, similarly, the starch in a potato before it can be used for the nutrition of the young shoots.

In the living organism these changes are brought about by the enzymes, and, in a word, enzyme action is the strategic centre of vital activity.

With regard to the mode of the formation of enzymes nothing is known; they are generally described as being due to the activity of the protoplasm, a phrase which contains no information. Sometimes the enzymes are secreted in specialized organs or in tissues more or less remote from the cells containing the material to be acted upon. In other cases they are formed in the same cells as the substrate.

A few examples may be given. In Zea Mais the cells of the surface of the scutellum next the endosperm have a distinct gland-like appearance, and here and there they dip down into the deeper layers of the scutellum, giving an appearance

^{*} Berzelius: "Jahresber.," 1850, 15, 237, 240, 278.

not unlike the crypts of Lieberkühn of the intestine. These secretory glands of the maize, however, have no lumina. In *Phanix dactylifera* the secretory organ of the seed is the rounded structure situated opposite the furrow. In *Nepenthes* and other insectivorous plants special glands occur in appropriate places, e.g. in the lining membrane of the pitchers, or in special tentacles, as in *Drosera*.

The fruits of *Rhamnus infectorius* are much used for dyeing. The pericarp contains a glucoside, xanthorhamnetin, which, on hydrolysis, breaks up into glucose and rhamnetin, a yellow compound. This hydrolysis is brought about in nature by an enzyme which is contained in the parenchyma of the raphe o the seed. To illustrate this, the following experiment may be tried.

An aqueous extract of the separated pericarp is made and placed in a glass vessel, then into the solution is thrown the raphe of a seed. A golden yellow precipitate comes down.*

In other cases the enzyme and substrate are contained in different cells of the same tissue, so that it is only necessary to crush up the tissue, or to macerate it, in order to obtain the reaction; the bitter almond, containing emulsin and amygdalin, may be given as an example.

The enzyme-secreting cells of Zea and Phanix have been studied by Reed.† He finds that in the resting condition these elements are crowded with granules of a protein naturel which disappear as secretion begins. At the beginning of secretion, the nucleus is poor in chromatin, but this material increases in amount as germination proceeds, the nucleolus becoming smaller and smaller. Finally, at the end of the secretory activity, the protoplasm of the gland-cells breaks down, and the products of its disintegration disappear from view.

It may be remarked that in the dried condition enzymes may retain their characteristic power for a considerable time; thus White; found that the ferments—diastase, protease and ereptase—of the resting fruits of wheat and barley retained

^{*} Ward and Dunlop: "Ann. Bot.," 1887, 1, 1.

⁺ Reed: id., 1904, 18, 267; see also Huie: "Q.J.M.S.," 1897, 39, 387; 1899, 42, 203.

[#] White: " Proc. Roy. Soc., Lond.," B., 1909, 81, 417.

their activity after twenty years, by which time the power of germination is lost. Also, that the subjection of the dry grains to certain extremes of temperature did not destroy the enzymes. Thus the heating of dry oats to 100° C, for four and a half hours was without effect in the destruction of the enzymes; an exposure to a temperature of 130° for one hour, however, did destroy the ferments. On the other hand, a temperature of - 200° C. did not destroy the dry diastase of barley.

The number of enzymes which a plant may contain is surprising; thus in Beta vulgaris, the leaves contain invertase, diastase, and maltase, the stem possesses invertase, diastase, inulase and emulsin, and the root diastase, maltase, inulase, and emulsin, but not invertase.*

The moulds—the digestive activities of which are, to a great extent, extra-cellular—also exhibit marked powers of secreting different enzymes. Thus Monilia sitophila may form maltoglucase, trehalase, raffinase, invertase, cytase, diastase, lipase, tyrosinase, and trypsin. These, according to Went,† are secreted according to the nature of the food; Dox, t however, who has demonstrated the presence in moulds of protease, nuclease, amidase, lipase, emulsin, amylase, inulase, raffinase, sucrase, maltase, lactase, histozyme, catalase and phytase. considers, from the data at hand, that these enzymes are formed regardless of the chemical nature of the substrate.

Observations such as these open up many questions relating to the nature of enzymes; are all these different ferments really specific, or are there only a few enzyme-nuclei which, before they can attack any particular substance, have to have attached to them certain molecular complexes according to the nature of the substrate?

There may, in certain cases, be made out a curious association of different enzymes. Thus Vines § found that when a tissue gave the guaiacum reaction, with or without the addition of peroxide, that same tissue also exhibited proteolytic activity and vice versa. Thus in the fruit of the orange, neither the juice nor the pulp gives the guaiacum reaction, whilst, on the

^{*} Robertson, Irvine, and Dobson: "Biochem. Journ.," 1909, 4, 258.

[†]Went: "Jahrb. Wiss. Bot.," 1901, 36, 611; see also Pringsheim and Zempter: "Zeit. physiol. Chem.," 1909, 62, 367.

‡Dox: "Plant World," 1912, 15, 40.

[§] Vines: "Ann. Bot.," 1903, 17, 257.

other hand, the peel does. The peel is actively proteolytic, but not the pulp and juice. Similarly the latex of the fig, papaw, lettuce, and spurge, has proteolytic qualities and also gives the peroxidase reaction. The meaning of this association is not clear.

CLASSIFICATION OF ENZYMES.

The following classification of enzymes, based on the chemical reactions in which they exert their accelerating influence, indicates the extensive use made by nature of these catalysts:—

I. HYDROLYTIC ENZYMES.

- (a) Ester or fat-splitting enzymes (esterases): Lipase.
- (b) Carbohydrate-splitting enzymes (carbohydrases):-

Invertase which hydrolyses cane sugar to dextrose and levulose.

,, ,, ,, raffinose to levulose and melibiose.

Maltase ,, maltose (malt-sugar) to dextrose.

Lactase ,, , lactose (milk-sugar) to dextrose and galactose.

Amylase or Diastase which hydrolyses starch to maltose and dextrin.

Inulase which hydrolyses inulin to levulose.

Pectinase * which hydrolyses pectose to arabinose.

Cytase which hydrolyses hemicellulose to mannose and galactose.

(c) Glucoside-splitting enzymes:-

Emulsin which hydrolyses amygdalin to glucose, hydrocyanic acid and benzaldehyde.

,, β-methylglucoside to glucose and methyl alcohol.

Myrosin ,, potassium myronate to allylisothiocyanate, glucose and potassium hydrogen sulphate.

Phytase ,, , phytin to inosite and phosphoric acid.

(d) Protein-splitting enzymes:— Pepsin contained in the stomach which hydrolyses proteins to

albumoses and peptones.

Trypsin , , pancreas which hydrolyses proteins to poly-

peptides and amino-acids.

Erepsin ,, intestine which hydrolyses proteins to poly-

peptides and amino-acids.

Bromelin ,, pine-apple juice which hydrolyses proteins to

Papain , , , pine-appie juice which hydrolyses proteins to polypeptides and amino-acids. , , juice of the fruit and leaves of the papaw

tree (Carica papaya) which hydrolyses proteins to polypeptides and amino-acids.

(e) Urea-splitting enzymes (ureases):-

Ureases obtained from *Micrococcus ureæ* and also from the Soja bean and other seeds † which hydrolyse urea into ammonia and carbon dioxide.

^{*} See p. 129. † Annett: "Biochem. Journ.," 1914, 8, 449.

- 2. FERMENTING ENZYMES.
 - (a) Alcoholic fermentation of glucose, levulose, mannose, etc., by zymase.
 - (b) Lactic acid fermentation of lactose by lactic acid bacteria.
 - (c) Butyric acid fermentation of lactose by the butyric bacteria, etc.
- 3 COAGULATING ENZYMES.

Rennin (Chymosin) which curdles milk.

Thrombin which coagulates blood.

Pectase .. soluble pectic bodies.

- 4. OXIDIZING ENZYMES.
 - (a) Oxidases which oxidize alcohols to acids, e.g., the action of Mycoderma aceti, etc., etc.
 - (b) Catalases or peroxidases which set free oxygen from hydrogen peroxide, or other peroxides, causing these substances to blue guatacum resin.

METHODS EMPLOYED IN ISOLATION OF ENZYMES.

The material from which the enzyme is to be extracted is ground up with water or dilute alcohol together with a little toluene to act as an antiseptic; sometimes the material to be extracted is previously dried by gently warming or by dipping in absolute alcohol; in some cases it is necessary to destroy the cell walls, before extraction, by grinding up with glass or Kieselguhr (infusorial earth) or by autolysis.

From aqueous solutions enzymes may be precipitated in the form of amorphous powders by the addition of an excess of alcohol.

Details for the isolation of certain enzymes are given below.

CHEMICAL CONSTITUTION.

The chemical constitution and nature of enzymes is, as yet, largely a matter of speculation, owing to the fact that it is very difficult to obtain an enzyme in a pure condition; attempts at purification generally end in the diminution or complete destruction of the activity of the material under examination. Owing to their tendency to be withdrawn (adsorbed) from solution by precipitates formed in their presence, it is difficult to purify them from proteins by any means which involve the precipitation of the latter; this may, to some extent, account for the fact that all enzymes were formerly supposed to be of a protein nature. According to Pekelharing * pepsin is in some way related to the nucleo-

^{*} Pekelharing: "Zeit. physiol. Chem.," 1885, Q. 577.

proteins although it contains no phosphorus; on the other hand, the purest forms of invertase hitherto obtained contain but little nitrogen and do not give the biuret reaction; they are rich, however, in carbohydrate and contain organically combined phosphorus.

Considerable difference of opinion exists in regard to the special class of enzyme known as oxidases. These, according to some authors, as for example Dony-Henault,* are not organic compounds at all, but owe their action to the presence of certain inorganic salts, more especially manganese salts, in colloidal solution. Bertrand, on the other hand, considers that the laccase of Rhus succedanca is a protein. whilst Euler and Bolin t are of the opinion that the laccase of Medicago sativa is composed of the calcium salts of glycollic, citric and malic acids.

According to Wolff, moreover, a very dilute ferrocyanide solution mixed with a colloidal iron solution gives all the reactions of an oxidase and is partly destroyed by boiling or mixture with traces of metallic salts.

PROPERTIES OF ENZYMES.

A peculiar property of enzymes, in which they differ from inorganic catalysts, is their sensitiveness to heat and light.

All enzymes are destroyed at 100° and most of them cannot, with safety, be heated much above 60°. The statement made by van't Hoff that the velocity of a reaction is doubled for every 10° rise of temperature is found to hold good for enzymes also with this reservation, that as the temperature approaches a certain height it begins to have a deleterious effect upon the enzyme; the so-called optimum temperature for the activity of any particular enzyme is therefore a compromise between the maximum acceleration effect, which can be attained in accordance with van't Hoff's rule, and the maximum temperature to which the enzyme can be heated without undergoing destruction. Inasmuch as the enzymes themselves are not living-unless, indeed, we consider the phenomena of life to be due to the activities of a complex of

^{*} Dony-Henault: "Bull. Acad. Roy. Belg.," 1908, 105. † Bertrand: "Ann. Chim. Phys.," 1907, [7], 12. ‡ Euler and Bolin: "Zeit. physiol. chem.," 1909, 61, 1. § Wolff: "Compt. rend.," 1908, 147, 745.

enzymes—the sensitiveness of an active enzyme, dissociated from the living cell, to heat is most readily explained by attributing it to the colloidal nature of the enzyme with the consequent tendency to coagulation by heat.

With regard to the action of light rays on enzymes it appears, according to Iodlbauer and v. Tappeiner,* that there exist two distinct kinds of action :-

(a) Those produced by ordinary light in the presence of oxygen, and (b) those produced by ultra-violet light independently of oxygen.

The destructive action which has resulted from exposure to bright sunlight therefore appears to be dependent on the presence of oxygen, and is greatly increased by the presence of fluorescent substances, such as eosin, quinoline red, etc.+

It was first shown by Green that ultra-violet light destroyed diastase, and since then several other authors have described similar effects for other enzymes.

The action of radium and radium emanation on enzymes has been studied by Wilcock, \$ by Loewenthal and Edelstein, by Bickel, by Loewenthal and Wohlgemut, and others.

The influence of various chemicals on the activity of enzymes will be dealt with later under the heading of " paralysers".

COLLOIDAL NATURE OF ENZYMES.

A fairly detailed account of the nature of colloidal solutions has been given above; it will suffice, therefore, merely to mention here that enzymes possess most of the more important properties of such solutions.

Foremost amongst these properties is their want of diffusibility; as already pointed out, this does not mean that they are quite unable to diffuse, but rather that their rate of diffusion is very small.** Their ability to diffuse through a membrane—commonly known as dialysis—is largely dependent on the nature or structure of the membrane, but, as a

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* Iodlbauer and v. Tappeiner: "Deut. Archiv Klin. Med.," 1906.
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⁺ Tappeiner: "Biochem. Zeit.," 1908, 8.

[#]Green: "Trans. Roy. Soc., Lond.," 1897, 188, 167.

^{\$} Wilcock: "Journ. Physiol." 1907, 34. || Loewenthal and Edelstein: "Bloch. Zeit.," 1908, 14, 484.

[¶] Loewenthal and Wohlgemut: "Biochem. Zeit.," 1909, 21, 476.

^{**} Chodiajew: "Arch. Phys.," 1808, 211.

general rule, enzymes will not pass through a parchment membrane; nevertheless Fränkel and Hamburg found that on subjecting a sample of diastase obtained from malt to dialysis, they were able to effect a separation into two distinct enzymes; one of these passed through the parchment and was found to be a sugar-producing enzyme, while the other which would not diffuse was able to liquefy starch.

On the other hand, most enzymes will pass through a porcelain filter, a fact which is made use of for separating the active enzymes from living cells. Owing to adsorption on the surface of the porcelain the filtration may, however, be accompanied by considerable loss of enzyme.

Enzymes for the most part are soluble in water, or in dilute salt solutions, or in glycerin. The lipases or fat-splitting enzymes, however, whether of animal or of vegetable origin, are characterized by their slight solubility in water.

On the other hand, enzymes are precipitated from solution by alcohol and by neutral salts such as ammonium sulphate.

Enzymes exhibit in a marked degree the phenomenon of adsorption,* and consequently are liable to be withdrawn out of solutions by other substances, such as calcium phosphate or uranyl phosphate, which may happen to be precipitated in their presence. For the same reason they are extracted from solution by shaking with charcoal, china clay, etc. The conditions obtaining here are to a large extent dependent on the electric charges of the substances concerned, a question which has been considered in detail by Michaelis.†

MODE OF ACTION OF ENZYMES.

To explain the mode of action of inorganic catalysts, it is frequently supposed that they form labile additive compounds with one of the reacting substances which then react more readily than the original substance would have done.

Similarly, in the case of the enzymes, it is now generally assumed that they enter into some form of loose combination with the substrate; in spite of this the enzyme is, in general, not altered by the reaction but retains its original activity

^{*}See Dauwe: "Hofm. Beitr.," 1905, 6, 426.

[†]Michaelis: "Biochem. Zeit.," 1908, 10, 283; "Dynamik d. Oberflächenwirkung," Leipzig, 1909.

after having completed its work, unless, of course, the products of the reaction have any effect on it.

In the group of carbohydrates the action of the enzymes is usually regarded as being more or less specific, each disaccharide being hydrolysed only by its own enzyme, e.g. cane sugar by invertase, milk sugar by lactase, and malt sugar by maltase.

That this specific activity is in some way connected with the molecular structure of the substances would appear from the researches of Fischer on the action of enzymes upon artificial glucosides. Fischer, by the action of methyl alcohol and hydrochloric acid on glucose, obtained two stereoisomeric methyl glucosides known respectively as the α and β variety. Now these two substances differ from each other only by the arrangement in space of the groups attached to the terminal carbon atom, and it is found that while the α modification is readily converted by maltase into glucose and methyl alcohol, the β modification is not affected by maltase at all, but is, on the other hand, hydrolysed by emulsin, which has no action on the α compound.

It would appear from this that the structure of the molecule which is to be decomposed is the determining factor.

Incidentally it may be mentioned that the fact that emulsin and maltase are complementary in their action upon a and β methyl glucosides, enables one to classify a glucoside as belonging to the a type if it is attacked by maltase and not by emulsin, or to the β type if it is attacked by emulsin and not by maltase.

Several other examples of this selective action on the part of enzymes for different optical isomers have been described by Fischer and Abderhalden, who found that whereas d-alanyl-d-alanine, d-alanyl-l-leucine were split up by enzymes, their stereoisomers d-alanyl-l-alanine and l-alanyl-d-alanine were not.

This peculiar dependence upon structure has led Fischer to suggest that the relationship which exists between the substance to be decomposed and its enzyme is similar to that existing between a lock and its key; or, in other words, unless the molecular structures of the two substances fit each other no interaction can take place.

These facts, of course, give strong support to the theory of

he formation of some sort of compound between the enzyme and the substrate

It should, however, be noted that the action of enzymes is not entirely specific, inasmuch as the one and the same enzyme may be able to hydrolyse two or more substances. Thus maltase is able to hydrolyse both maltose and a methyl glucoside; and emulsin is able to decompose β methyl glucoside, β methyl galactoside, milk sugar, amygdalin (the glucoside of bitter almonds, and with which it is primarily associated in nature), arbutin, salicin, and coniferin.

The specific nature of the interaction between enzymes and other substances is, however, only really strongly marked in connexion with optically active substances. For, taking the case of the fat-splitting enzymes or lipases, practically all esters are broken up by pancreatic lipase, although the ease with which the hydrolysis is effected may vary considerably in different cases.

On the other hand, Fischer and Abderhalden have shown that whereas pancreas extract was able to hydrolyse a number of artificial polypeptides, it was quite unable to act upon others.

Fischer has described enzymes as optically active catalysers, and explains in this way how it is that they produce optically active substances from inactive material, as, for example, when moulds, such as *Penicillium glaucum*, *Aspergillus niger*, etc., are allowed to grow upon inactive tartaric acid with the formation of l-tartaric acid; it is assumed that the enzymes combine with the racemic substrate to form isomeric substances which decompose at different rates and so form optically active products.

A most important contribution to the elucidation of the activity of enzymes is the discovery of the stimulating or inhibiting influence exercised by certain substances which may be described respectively as Activators and Paralysers.

ACTIVATORS.

Under this heading are included substances described by different authors as Co-enzymes * or Accelerators. In some

^{*}Bayliss distinguishes between co-enzymes and accelerators by reserving the former term for those substances without which the enzyme is unable to

cases the substances are quite simple chemical individuals, such as acids, alkalis, or salts, and in others they may be complex and of unknown constitution, as in the case of the co-enzyme of zymase (but see p. 378). They have, however, properties in common, namely that they can be separated from the enzymes by dialysis, and are not destroyed by heating. Moreover, an enzyme rendered inactive by removal of its co-enzyme can be restored to its original activity by mixing again with this substance.

This latter effect has been shown especially for dialysed liver extract which has no lipolytic action; if, however, this extract be mixed with some of the solution which had been dialysed out, or with boiled liver extract, the characteristic lipolytic action is regained. In this case it can be shown that the bile salts are the active co-enzyme. Similarly, the activity of zymase, the enzyme of yeast cells, is dependent on the presence of certain complex phosphoric esters, which, likewise, can be separated from zymase by dialysis and are not destroyed by boiling water.

Other examples of the dependence of the enzyme upon activators are the necessity for the presence of a small quantity of acid in order that pepsin and the lipase of castor-oil seeds, for instance, may exert their respective actions.

Similarly trypsin requires a faintly alkaline medium to exert its proteoclastic action; in many cases the presence of calcium salts is essential, as, for example, in the clotting of milk by rennin, the clotting of blood by thrombin, and the gelatinization of pectin by pectase.

In the case of some enzymes the substance at its seat of origin is not a true enzyme but a so-called proferment or zymogen which is not itself active but only becomes so on being brought into contact with another more or less complex substance known as a kinase. Thus, for example, trypsinogen, which is contained in pancreatic juice, has very little action on proteins but is converted into the true proteolytic enzyme trypsin on coming in contact with the kinase—entero-kinase

exert its activity, and the latter for substances which stimulate or accelerate a reaction without being absolutely essential to its taking place; an example of the latter class is furnished by traces of manganese salts which greatly increase the oxidizing power of the enzyme laccase, though it has yet to be proved that the laccase is unable to act in their absence.

-which is secreted by the mucous membrane of the duo-denum.

The relation between proferment and kinase is different from that existing between enzyme and co-enzyme inasmuch as the two latter can be alternately mixed and separated; on the other hand, the reaction between proferment and kinase is not reversible; furthermore, the proferment is not really regarded as a complete ferment, while the true ferment produced from it, by combination with the kinase, may still be dependent upon an activator for its activity.

The following example of the dependence of thrombin upon calcium salts will illustrate this; the coagulation of blood by thrombin consists in the conversion of the soluble substance fibrinogen into the insoluble substance fibrin. The blood plasma contains a proferment thrombogen and also calcium salts, but these two substances alone are unable to coagulate fibrinogen. When, however, the blood is drawn, a kinase, known as thrombokinase, which is secreted by the blood corpuscles, combines with the proferment thrombogen forming the true coagulating ferment thrombase.

Both enterokinase and thrombokinase are destroyed by heat.

PARALYSERS.

Paralysers are substances which reduce or destroy the activity of enzymes. These may be either the products of the activity of the enzyme or of foreign substances. Examples of the first class are the acetic or lactic acids which, unless neutralized, destroy the ferments producing them; similarly the alcohol produced by the fermentation of sugar ultimately stops the fermentation. Also, Tammann has shown that the hydrolysis of amygdalin by emulsin was retarded by the addition of any one of the products of hydrolysis, namely glucose, benzaldehyde or hydrocyanic acid, but most markedly by the latter. Similarly Croft-Hill found that glucose interfered with the action of maltase, and the Armstrongs* likewise have pointed out a number of examples of the inhibiting action of the reaction-products upon the enzyme.

Amongst foreign substances having a retarding effect on

^{*} Armstrong: "Proc. Roy. Soc., Lond.," B., 1907, 79, 360.

enzymes may be mentioned inorganic substances such as mercuric chloride or cyanide, arsenious oxide, sulphuretted hydrogen, ozone, and organic compounds such as chloroform, chloral, formaldehyde, hydrocyanic acid, phenylhydrazine aniline, alcohol, etc.; the influence of these substances on different enzymes varies considerably; thus, for example, alcohol usually acts as a paralyser, but on lipase it has a stimulating effect.

The majority of the substances included in the above list also act as poisons to colloidal solutions of metals; the peculiar phenomenon of the recovery of metallic colloidal solutions from poisoning by hydrocyanic acid, is also met with in the case of the enzymes, and is likewise attributed to the oxidation of the poison.

The mechanism of toxic action is as yet unexplained; it is assumed that some form of chemical combination between the paralyser and the substrate enzyme or activator takes place.*

The work of Caldwell † on the effect of toxic agents upon bromelin, the proteolytic enzyme of the pineapple fruit, may be cited in illustration. The prepared enzyme (see p. 370) was dissolved in water so that each cubic centimeter contained '006 gram and the solution was rendered acid or alkaline by the addition of hydrochloric acid or sodium hydrate of a concentration of M/32. Five c.c. of the enzyme solution was placed in a series of test tubes together with I gram of boiled granulated egg albumen; then to each tube was added a solution of the paralyser.

The tubes were then placed in a water bath and kept at a temperature of 40° C. for twenty-four hours. At the end of this period, the liquids were filtered, if necessary, to remove any albumen, and tested for peptones, leucine and tyrosine by the biuret and tryptophane reactions, confirmatory tests being applied if necessary.

In the following table the metals are arranged in their order of toxicity, the top ones being the most poisonous, and are compared with the results obtained by Matthews !

^{*} Cf. Loewenhart and Kastle: "Amer. Journ. Chem.," 1903, 29, 397, 563.

[†] Caldwell: "Bot. Gaz.," 1905, 39, 409. ‡ Matthews: "Ann. Journ. Physiol.," 1904, 10, 290; 1904, 11, 455.

on the eggs of Fundulus heteroclitus and McGuigan* with diastase

Matthews.	McGuigan.	Caldwell
Ag	Ag	Ag
Hg	Hg	Hg
Cu	Cu	Cu
Cd	Cd	Pb
Pb	Co	Zn
Zn	Zn	Ba
Co	Pb	Cd
Li	Sr	Co
Sr Na	Ba	Na
	Mg	Li
Ba	Li	Sr
Mg	Na	Mg
NĤ₄ K		NH₄ K

With regard to salts, Caldwell found that nitrates inhibit the action of the enzyme in somewhat greater dilution than the corresponding sulphates and chlorides. He agrees with Matthews that the affinity of the atom or ion for its electrical charge is the main factor which determines its physiological action.

It is to be remembered that the effect of poisons vary with the purity of the preparations used; a slight admixture of proteins and other impurities makes it necessary to increase the concentration of the poison greatly in order to inhibit the enzyme action.

ANTI-ENZYMES.

The term anti-enzyme is applied to a class of substances occurring in the living organism or produced in it by subcutaneous injection with an enzyme. The anti-enzymes are antagonistic in their action upon the enzymes, and their action is quite specific, the relationship between an enzyme and its anti-body being similar to that existing between a toxin and an anti-toxin. The first example of immunity against an enzyme was recorded by Hıldebrandt,† the enzyme being emulsin.

^{*} McGuigan: id., 1904, 10, 441.

⁺ Hildebrandt : "Virch. Arch.," 1893, 131, 12, 26.

Since then, anti-enzymes have been discovered for lipase, amylase, pepsin, papain and urease. Anti-trypsin and anti-rennet occur normally in the blood, and, according to Weinland,* anti-pepsin and anti-trypsin occur in the mucous membranes of the stomach and intestine respectively.

In this connexion the work of Czapek † on the anti-ferment reaction in tropistic movements of plants is of particular interest. It is impossible here to give a complete account of the investigations referred to, but the following facts will give some idea of the phenomena under discussion.

- I. It was found that the roots of *Vicia Faba* when subjected to the stimulus of gravity always reduced silver more effectively than unstimulated roots. The mode of testing is as follows. The roots, stimulated or otherwise, are cut into longitudinal slices and boiled in an ammoniacal solution of silver nitrate; the darkness assumed by the preparations is an indication of the amount of silver reduced.
- 2. If the roots of the lupin, for example, be anæsthetized, a deposition of tyrosine, in the shape of spherical crystals, takes place; but, strangely enough, not in the root-tip, nor in the youngest parts of the growing regions where the maximum reduction of silver takes place.

The question naturally arises, is there any connexion between the reduction of silver and the production of tyrosine? As a matter of fact there is, for it was observed that in roots containing much crystalline tyrosine, although at first the tyrosine-containing cells did not reduce silver, they did so eventually, the silver reducing properties becoming more and more marked as the tyrosine disappeared, so that eventually a very strong reduction obtained.

By these and similar observations it was shown that the tyrosine disappears by means of a ferment, tyrosinase, and that one of its products is the substance which reduces the silver. Tyrosinase appears to be widely distributed in plants, e.g. it has been identified in *Russula*, in the tubers of the *Dahlia*, in the beetroot, as well as in root-tips. The product of the decomposition of tyrosine referred to above is homo-

^{*} Weinland: "Zeit. f. Biol.," 1903, 44, 45. + Czapek: "Ann. Bot.," 1905, 19, 75.

gentisinic acid (C₈H₈O₄),* a substance which is also produced in a similar fashion in the human body.+

HO
$$CH_2CHNH_2COOH + 3O = OH$$
 $CH_2COOH + CO_2 + NH_3$

Homogentisinic acid

Homogentisinic acid may be prepared from root-tips by grinding them in 96 per cent alcohol, filtering off the solids, and evaporating over a water bath. The residue when dissolved in water forms a brown solution which is free from sugar, and gives a faintly acid reaction. In the air it turns dark.

The following are characteristic reactions.

- Treated with alkali it turns reddish-brown.
- 2. It reduces an ammoniacal solution of silver nitrate on warming.
 - 3. Fehling's solution is feebly reduced by it on warming.
 - 4. Ferric chloride gives a green coloration.
 - 5. Ferric sulphate gives a violet-blue coloration.
 - 6. Millon's reagent gives a yellow colour.
 - 7. Homogentisinic acid is precipitated by lead acetate.
 - 8. It also gives a reddish colour with hydrogen peroxide.

Czapek further found that if ground-up root-tips were kept for some time under the influence of chloroform, the mass gradually loses its power of reducing silver, and even small quantities of homogentisinic acid intentionally added to the preparation gradually disappear; this is due to the action of an oxidase.

It has been seen that root-tips stimulated by gravity give a strong reduction of silver, but not so the unstimulated roots. To explain this there are two alternative hypotheses; either the action of the oxidase is inhibited during geotropic stimulus, so that the homogentisinic acid, which is otherwise acted upon. disappears more slowly than under ordinary circumstances and so accumulates, or, there is a diminution in the production of oxidase by the root-tip.

+ Wolkow and Baumann: id., 1891, 15, 260; Huppert: "Z. physiol.

Chem.," 1897, 23, 412; Garrod and Hele: id., 1905, 33, 198.

^{*} The occurrence of this acid in plants is denied by Schultze and Castoro: "Z. physiol. Chem.," 1906, 48, 387, 396. See also Bertel: "Ber. deut. bot. Gesells.," 1902, 20, 454.

From experimental evidence, Czapek considers that the first alternative is the true one. He found reason to believe that the inhibiting substance is an anti-oxidase of some potency which is precipitated by alcohol, destroyed by heat (62°) and may be isolated by filtration through a porcelain candle. Czapek further ascertained that the anti-oxidase is more or less specific, for although the oxidase and the anti-oxidase of closely related plants have a mutual action, this is not so for plants widely separated.

It is supposed that the anti-oxidase is only produced as a result of gravitation stimulus, so that the simple reaction is as follows. The tyrosine is converted by the action of tyrosinase into homogentisinic acid; the further oxidation of the acid by oxidase is inhibited by the production of an anti-oxidase which renders the oxidase more or less inefficient, and so the homogentisinic acid accumulates.

For the mode of quantitatively determining the amount of homogentismic acid and for other details, Czapek's paper must be consulted.

The interaction between enzyme and antienzyme appears to be of the nature of adsorption.

ENZYMES AND THE LAWS OF MASS ACTION.

According to the Law of Mass Action enunciated by Guldberg and Waage, the rate at which a body undergoes chemical change is dependent on the concentration as measured by the number of gram molecules of substance present in the litre; consequently the amount of substance changed in unit time will be greater at the beginning of the reaction than towards the end, since the amount of unchanged substance is continually decreasing.

The relationship between the amount of substance x (measured in gram molecules per litre) changed in time t (measured in minutes) and the original concentration a of the substance is given by the equation:—

$$K = \frac{1}{t} \log \frac{a}{(a-x)}$$

The above formula holds only for the decomposition of a single substance, and it is, therefore, characteristic of what is known as a Monomolecular reaction or a reaction of the first order, and as such is applicable to all cases of hydrolysis, as for example:—

$$C_{12}H_{22}O_{11} \ + H_2O = 2C_6H_{12}O_6$$

Although from the left-hand side of the equation it would appear that two substances are reacting, the quantity of water present is so large, as compared with the amount of cane sugar, that its concentration is practically unaltered, and therefore, for all intents and purposes, only a single substance is undergoing alteration in concentration.

Now the hydrolysis of cane sugar which takes place slowly in aqueous solution is catalytically accelerated by the addition of dilute mineral acids, the effect being greater in proportion to the amount of acid used, without, however, altering the order of the reaction; in just the same way enzymes accelerate hydrolyses in accordance with the law of mass action for monomolecular reactions, thereby showing that they are true catalysts.

In reactions acting in accordance with the logarithmic equation above given, the amount of substance changed in a given time bears a constant ratio to, or is a constant fraction of, the amount of substance unchanged; on plotting the amounts changed as ordinates against the time as abscissæ there is accordingly obtained what is known as a logarithmic curve

Now it is found that when this is done for an enzyme reaction the curve both at the beginning and at the end of a reaction is not logarithmic but linear. Thus Horace Brown and Glendinning* found that equal amounts of starch were hydrolysed by diastase in equal times during the earlier part of the reaction, in other words, the course of the reaction was expressed by a straight line; as the reaction proceeded, however, it became logarithmic, or, in other words, at the commencement, when the concentration of the substance being hydrolysed is great as compared with that of the enzyme, the reaction is linear and not in accordance with the law of mass action, but where the concentration of the enzyme is great as compared with that of the substance being hydrolysed, the reaction obeys the law of mass action.

^{*} Brown and Glendinning: "J. Chem. Soc., Lond.," 1902, 81, 392.

Similar results were obtained by Adrian Brown* in the study of the action of invertase on cane sugar; he also expresses the view that, in the case of alcoholic fermentation and other enzyme actions which do not apparently conform with the law of mass action, the exceptional action "is due to a time factor accompanying molecular combination and change which limits the influence of mass action . . . this theory demands not only the formation of a molecular compound of enzyme and reacting substance, but the existence of this molecular compound for an interval of time previous to final disruption and change".

Similarly E. F. Armstrong † in studying the action of lactase and maltase upon their respective sugars found that while the reaction was in the main logarithmic, both the initial and final stages were linear; this is explained by the fact that as a result of the combination between the enzyme and the substrate there will be an excess of substrate at the commencement but an excess of enzyme at the end, both of which conditions favour a linear change.

On calculating the velocity constant for that part of the reaction which is logarithmic it is found that, as a rule, the value steadily decreases, or, in other words, the enzyme appears to become less active. This may be accounted for in one of two ways: either by the assumption that the products of the reaction combine with the enzyme or, by their concentration, exercise some inhibiting influence upon the enzyme; or else by assuming that the tendency for the reverse action to take place has a retarding effect.

That there should be a tendency for the reverse reaction to take place is a perfectly legitimate conclusion; in fact van't Hoff long ago pointed out that a catalyst which accelerates a reaction in one direction must also be able to exert an accelerating effect on the reverse reaction. Consequently the same enzymes which effect hydrolyses should also, under suitable conditions, be able to synthesize.

The first experimental proof of this was given by Croft Hill, ‡ who showed that when maltase was allowed to act on

^{*} Adrian Brown: "J. Chem. Soc., Lond.," 1902, 81, 379.

[†] Armstrong: "Proc. Roy. Soc., Lond.,"B., 1904, 73, 500, 516, 526; 74, 188, 195. ‡ Croft Hill: "J. Chem. Soc. Lond.," 1898, 73, 634.

a concentrated solution of glucose, the disaccharide iso-maltose was produced; later it was shown that the disaccharide iso-lactose could be synthesized from galactose and glucose by the action of lactase from Kefir. Since then a large number of enzymatic syntheses have been effected, amongst them being included the synthesis of maltose itself by the action of emulsin on a concentrated solution of glucose which was described by E. F. Armstrong, and also the formation of glycogen from a 30 per cent solution of fructose by yeast-extract free from glycogen,* a reaction which most probably involves the conversion of fructose into glucose.

The following experiment has been devised by Bayliss† for demonstrating the synthetic action of emulsin in the production of a glucoside:—

Two solutions are required: (a) a 15 per cent solution of hydroquinone in glycerol, and (b) a 50 per cent aqueous solution of glucose mixed with an equal volume of glycerol. The two solutions (a) and (b) are mixed in equal proportions, and about 2 per cent of emulsin are added, and the mixture is ground in a mortar and then warmed to 38° to make it less viscid. Ten per cent of the solution are then delivered by means of a pipette into a series of test tubes, a little toluene is added to each, and after displacing the air above the liquid by CO_2 , the tubes are sealed up in a blow-pipe, but this is not absolutely necessary; if this is not done the liquid may darken owing to oxidation of the hydroquinone, but on adding a small quantity of sodium bisulphite the colour is discharged.

One sample is at once diluted to 50 c.c., and filtered, and its rotation is measured in the polarimeter. The other tubes are placed horizontally in an incubator, and samples are withdrawn every three or four days; the tubes are cracked across, and their contents are rinsed into a 50 c.c. flask.

The initial rotation of about + 3° will be reduced to 0.5° in a week, indicating a synthesis of 25 to 30 per cent of arbutin.

To prove that the diminution of rotation is not due to destruction of glucose the glucoside may be reconverted into

^{*}Cremer: "Ber. deut. chem. Gesells.," 1899, 32, 2062. See also Meyer: "Bot. Ztg.," 1899, 57, 313. †Bayliss: "J. Physiol.," 1912, 43, Proceedings, xL.

its components by diluting a sample to 50 c.c. and adding half a gram of fresh emulsin. In about two to three days the hydrolysis will be complete, and the original rotation will be restored. As commercial emulsin is, however, itself levorotatory, a control experiment should be made with emulsin alone or else the emulsin may be precipitated by adding mercuric nitrate, and filtering before the polarimeter reading is made.

In a subsequent investigation * Bayliss found that actually very little arbutin was formed, but that the glucoside really produced was that of glycerol. That being so, the hydroquinone can be left out of the mixture, and as a result the sealing of the tubes and subsequent treatment with sodium bisulphite is rendered unnecessary. The best mixture is made as follows: Glucose (anhydrous) 18, water 12, glycerol 40, and emulsin 3 parts by weight. The glucose must be dissolved in the water and cooled before adding the glycerol, owing to the production of glycerol glucoside by heat. At a temperature of 47° a diminution of rotation from + 2°·83 to + 0°·80 takes place in seven days, and equilibrium at – 0°·16 (corresponding to about 75 per cent synthesis) is practically attained in fifteen days.

The experiment is of interest with regard to the theory of catalysts, since if the above mixture of glycerol, hydroquinone and glucose is heated without enzyme to 100° for some hours a certain amount of glucoside synthesis results; it may, therefore, be assumed that some synthesis also takes place, though very much more slowly, at 38°. The emulsin, therefore, in accordance with Ostwald's definition of an enzyme, merely accelerates a reaction which is already taking place, though very slowly.

A CONSIDERATION OF CERTAIN TYPES OF ENZYMES.

Before passing on to a more special consideration of individual enzymes, attention must be drawn to a point of general application to all enzymes. It is of the highest importance that experiments on enzymes which take several hours to carry out, should be conducted under aseptic conditions in order to avoid bacterial activity. The fermenting mixtures obviously cannot

^{*} Bayliss: "J. Physiol.," 1912, 44, Proceedings, ix, and 1913, 46, 236.

be sterilized by means of heat, so that antiseptics must be added. Amongst those commonly employed are chloroform, toluene, thymol, sodium fluoride, and hydrocyanic acid. The nature of the antiseptic exerts a considerable influence upon the activity of the enzyme used, so that it is necessary to try many different antiseptics. The following table illustrates this in the case of papain*:—

Reaction.	Hydrocyanic acid.	Chloroform.	Sodium fluoride.
Acid (·5 per cent citric acid)	Fibrin quite disintegrated Marked trypto- phane reaction	Scarcely attacked Faint trypto- phane reaction	Distinctly attacked Faint trypto- phane reaction
Alkaline (5 per cent Na ₂ CO ₃)	Fibrin quite disintegrated Faint trypto- phane reaction	Scarcely attacked Doubtful trypto- phane reaction	Scarcely attacked Doubtful trypto- phane reaction
Neutral	Fibrin nearly all gone Distinct trypto- phane reaction	Distinctly attacked Faint trypto- phane reaction	Mostly disintegrated Faint trypto- phane reaction

LIPASE.

The existence of a fat-splitting enzyme or lipase in the animal kingdom has long been known. This substance, which is known as steapsin, is contained in the pancreas; acting in an alkaline medium it is able to break up fats into glycerol and free fatty acids, the latter combining in the intestine with alkali to form the sodium salts or soaps.

In 1890 Green † found that germinating seeds containing fat or oil, when macerated with water and left for some time, gradually acquired an acid reaction. This observation was subsequently confirmed and extended by Connstein, Hoyer, and Wartenberg, ‡ with the result that it has been found that

^{*} Vines: "Ann. Bot.," 1903, 17, 602.

⁺ Green: "Proc. Roy. Soc., Lond.," 1890, 48, 375.

[‡] Connstein, Hoyer and Wartenberg: "Ber. deut. chem. Gesells.," 1902, 35, 3988; Hoyer: id., 1904, 37, 1444; "Zeit. Physiol. Chem.," 1907, 50, 414.

the seeds of Euphorbiaceæ, and especially castor-oil seeds, whether germinating or not, contain an enzyme capable of hydrolysing not only the fat present in such seeds but also fats from other sources. The observation that the hydrolysis takes place slowly at first and then suddenly increases from 5 per cent after one day to 58 per cent after two days and to 95 per cent after four days led the authors to the conclusion that for rapid hydrolysis a certain minimum amount of free acid must be present, and it was found that when a little free acid was added from the commencement hydrolysis could be completed within a few hours. Similar observations regarding the curve of the hydrolysis of fats during the germination of *Ricinus* seeds have been made by Delcano.*

According to Nicloux † fats may be attacked by other means. He used castor-oil seeds, which were ground up and the cytoplasm separated from the aleurone grains and other cell contents by mechanical means.

It was found that the cytoplasm thus prepared showed a marked power of hydrolysing fats, acting in the same way as an enzyme and obeying the laws of enzyme action. But inasmuch as the active substance, which Nicloux calls lipaseïdine, contained in the protoplasm is destroyed by water as soon as its protective layer of fat is removed, it is not considered to be an enzyme in the ordinary sense of the term.

The most favourable conditions for the activity of lipase may be summarized as follows:—

- (a) The presence of free acid varying from N/60 to N/100 or less, according to the amount of material to be hydrolysed.
 - (b) The presence of a certain amount of water.
 - (c) The formation of a good emulsion.
- (d) The maintenance of a suitable optimum temperature, varying from about 23° to 42° C.

THE ISOLATION OF LIPASE.

Lipase may be separated from the seeds of *Ricinus* by the following means: The seeds are allowed to begin germination; when the radicles have protruded a little way, the endosperms are ground in a mortar with a 5 per cent solution of sodium

^{*} Delcano: "Centrlbl. Bakt.," 1909, 24, 130.

⁺ Nicloux: "Proc. Roy. Soc., Lond.," B., 1906, 77, 454.

chloride. The liquid is filtered off and placed in a dialyser; the salt having thus been removed, an excess of alcohol is added and the precipitate filtered off. The precipitate is washed with alcohol and may be dissolved in water before use.

For commercial purposes the enzyme is prepared as follows:* Castor-oil seeds are ground up with water and then centrifuged; the resulting emulsion, which contains castor oil, proteins, and the enzyme, is then allowed to ferment at a temperature of 24°, whereby a scum containing the ferment rises to the surface and can be separated from the aqueous layer. This scum is then allowed to act upon the molten fat in the presence of water and a little manganese sulphate as a catalytic agent.

The following experiments described by Connstein, Hoyer, and Wartenberg, may be taken as an illustration of the process on a small scale

Five grams of castor-oil seeds are macerated with 10 c.c. of water containing 0.2 gram of acetic acid and 0.1 gram of chloral hydrate. After twenty-four hours it is found that about 58 per cent of the fat originally present has been hydrolysed.

To show that the enzyme is not destroyed by extracting the fat from the seeds by means of ether, 1.5 grams of seeds which had been so extracted were ground up with 75 grams of cotton-seed oil and 15 grams of N/10 sulphuric acid. In forty-four hours 82 per cent of the oil had been hydrolysed.

QUANTITATIVE DETERMINATION OF THE ACTIVITY OF LIPASE.

The activity of this enzyme is conveniently studied by allowing it to act on ethyl butyrate and observing the amount of acid liberated by titration or by conductivity measurements.

DIASTASE.

Diastase is one of the commonest of enzymes; in fact it may be regarded as being universally present in the higher plants. The amount present in any particular organ varies according to the conditions obtaining; thus when the temperature and other factors are most favourable for growth and for the germination of starchy seeds, diastase is much more abundant than when growth and germination are sluggish.

^{*} Cf. Hoyer: "Der Seifenfabrikant," 1905, 25, No. 27.

Also, the amount of diastase is always greater in starch leaves than in sugar leaves and the same holds for insolated leaves containing much starch, as compared with shaded leaves containing little or no starch.*

ISOLATION OF DIASTASE.

To obtain a relatively large quantity of diastase, germinated barley gives excellent results. The grains are soaked in water for twelve hours, and then spread out in a thin layer on a tray which is placed in a warm, damp—but not too damp—place. When the radicles are about one quarter of an inch long, the grains may be dried at a temperature not exceeding 40° C.; they are then ground up as finely as possible. The powder is mixed thoroughly with about four times as much water, and allowed to stand for an hour or two, the mixture being well shaken up periodically. The fluid is then filtered off and evaporated in a vacuum to a small bulk; this concentrated solution is poured into an excess of absolute alcohol, whereby the diastase, and other substances, are precipitated. precipitate is filtered off, and washed with alcohol. diastase thus obtained may be partly purified by dissolving in water and re-precipitating with alcohol.

Although diastase occurs in green leaves, it is often difficult to demonstrate its presence in an aqueous extract of the fresh tissue. If the leaves be dried and ground to a very fine powder, the above procedure should yield positive results; if not, then the powdered leaves may be added directly to a one per cent solution of starch paste or to a little dry starch suspended in water in a watch glass. The disappearance of the starch, as indicated by the iodine reaction, and the corrosion of the solid starch grains, point to the presence of diastase.

The action of this enzyme is promoted by the presence of acids, e.g., hydrochloric or citric, but if too much acid be added, the action is inhibited.

To study the action of diastase on starch a mixture of these two substances may be tested from time to time with indine solution.

^{*} Eisenberg: "Flora," 1907, 97, 347.

Appleman* gives the following experiment. A number of test tubes, say ten, each containing I c.c. of a one per cent solution of starch paste, are placed in ice. The extract of the material to be examined for its diastatic activity is added to the mixture in increasing amounts. Thus to the first tube is added I c.c. of extract, to the second I I c.c., to the third I 2 c.c., and so on. A drop or two of toluol are also added as an antiseptic. The tubes are then removed from the ice and placed in an incubator, kept at a temperature of 40° C., for forty-eight hours. An equal amount of water, roughly enough to fill the test tubes, is added to each test tube and, after shaking up, three drops of iodine solution. The first tube in the descending series which showed a blue or violet colour was taken as the index for comparison.

QUANTITATIVE DETERMINATION OF THE ACTIVITY OF DIASTASE.

The amount of optically active or reducing sugars produced may be followed polarimetrically or by means of Fehling's solution. Due allowance should be made for any sugar contained in the enzyme.

PROTEASES

For many years it has been known that the fluids excreted by many insectivorous plants are capable of digesting proteins; proteolytic ferments are now known to occur in the juice of a good many plants. Some indeed, e.g., erepsin, are almost universal. Amongst the better known ones may be mentioned papaïn which occurs in the fruit of Carica papaya (papaw), bromelin in the fruit of the Ananas sativa (pineapple), and cradeïn in the latex and fruit of Ficus (fig).

ISOLATION OF THE ENZYME.

The methods followed in isolating these enzymes differ in details according to the material used; the principle, however, is the same in most cases. The enzyme is precipitated from its solution by reagents, usually alcohol, filtered off, and washed with alcohol. It may be partly purified by dissolving in water and re-precipitating with alcohol. Following are some methods which have been pursued in particular cases.

^{*} Appleman: "Bot. Gaz.," 1911, 52, 306.

To isolate the enzymes from the fluid contained within the pitchers of Nepenthes, Vines * added to the liquid an equal volume of absolute alcohol, then phosphoric acid followed by lime water in order to increase the bulk of the precipitate. Ammonium carbonate was added until the liquid gave a neutral reaction, and the precipitate filtered off. For use, the precipitate was shaken up with a '2 per cent solution of hydrochloric acid and filtered; the clear filtrate actively digests fibrin.

If it be desired to examine the contents of a tissue for these ferments, the expressed juice may be used, or an aqueous extract, the enzyme being separated as above if necessary. But sometimes this is unsatisfactory for various reasons—a syrup-like consistency or high coloration, for example. such cases the tissues may be bruised in a mortar and placed with water in the vessel in which the experiment is to be carried out, together with the material-fibrin, for exampleto be acted upon.+ Buscalioni and Fermi! used sterilized gelatine, with 5-1 per cent carbolic acid as an antiseptic, in a Petri dish. Fragments of the tissue to be tested are placed upon the jelly; the liquefaction of the gelatine in the neighbourhood of the pieces indicates the presence of proteolytic enzymes, but inasmuch as all proteases do not attack gelatine. a negative result does not necessarily indicate the absence of these enzymes.

Dean \$ prepared ereptase from the seeds of beans by extracting the cotyledons with water, filtering, and half saturating the filtrate with ammonium sulphate. The precipitate thus obtained is filtered off, dissolved in water and separated from ammonium sulphate by dialysis. The solution of enzyme thus purified may be dried at a temperature below 50° C.

Vines || separated peptase from ereptase by making use of the fact that the former is hardly soluble in water but readily so in a dilute solution of sodium chloride, whilst ereptase is easily soluble in water. The material, e.g., seed of *Cannabis sativa*, is ground and extracted with a 10 per cent solution of

^{*} Vines: "Ann. Bot.," 1897, 11, 573. † Vines: id., 1903, 17, 237, 597. ‡ Buscalioni and Fermi: "Ann. R. Inst. Bot., Roma," 1898, 7, 99. § Dean: "Bot. Gaz.," 1905, 39, 321. || Vines: "Ann. Bot.," 1908, 22, 103.

sodium chloride. The solution is filtered and rendered just acid by the addition of acetic acid, whereby a white precipitate of protein is formed, which is filtered off. The acid filtrate has marked proteolytic qualities but has no action on fibrin; it therefore contains the ereptase. The fibrin-digesting protease (peptase) is in the precipitate; to recover it, wash the precipitate with a 10 per cent solution of sodium chloride slightly acidified with acetic acid. The precipitate is next treated with distilled water and filtered; the filtrate, which has an opalescent appearance, digests fibrin but has no effect on Witte peptone. In order to ensure the best results, the temperature should be kept as low as possible during filtration.

GENERAL CONSIDERATIONS.

According to Vines, the proteases of plants fall into two main groups:—

- I. Peptase.
- 2. Ereptase.

Peptase hydrolyses proteins to albumose or peptone, but does not act on albumose or peptone whether produced by its own digestion of protein or added in the form of Witte peptone.*

Ereptase hydrolyses proteins, albumoses and peptones to amino acids, such as leucine and tyrosine.

Peptase dissolves readily in a saline solution but is hardly soluble in water, whilst ereptase is easily soluble in water. Both may occur in a plant, e.g. the seeds of *Cannabis sativa*,† the latex of *Carica papapa*—the enzyme of which is termed papan—yeast, etc.‡ In fact the mixture is, or was, commonly termed vegetable trypsin.

On the other hand, some plants which exhibit proteoclastic properties only have peptase. This is, however, seemingly very rare; *Drosera* provides an example.§

Although ereptases are very common in plants, peptases are less common, and have not been found in some cases where they might be expected to obtain, e.g. in protein-containing seeds. Dean's work on *Phaseolus vulgaris* may be taken as an

^{*} Vines: "Ann. Bot.," 1905, 19, 171; 1908, 22, 103.

⁺ Vines: id., 1908, 22, 103.

[‡]Vines: id., 1909, 23, 1.

[§] White: "Proc. Roy. Soc., Lond.," B., 1910, 83, 134.

example.* The seeds of this plant contain much protein which undergoes proteolysis before translocation takes place. But no enzyme has been discovered in the seed which is capable of digesting these proteins; ereptase, however, which can hydrolyse the proteases derived from the digestion of these seed proteins, is abundant. Dean considers that the protoplasm plays the part of a peptase, whilst the ereptase may carry the digestion further.

The plant proteases are less rigid than the corresponding ones from animal sources in respect to their activity in acid or alkaline media. Thus the proteolytic enzyme of *Drosera* is active in acid, alkaline or neutral media; papaïn is active both in acid and alkaline media, thus differing from animal pepsin;† and some proteases will only work provided the reaction be acid, e.g., *Nepenthes*, malt, mushroom and yeast.‡ The natural reaction of the plant juice is the best to maintain for general experiments.

TRYPTOPHANE REACTION.

The presence of tryptophane is an indication of the activity of trypsin-like proteolytic ferments. Tryptophane may occur naturally in the sap of the plant, its presence being associated with the ripening of fruits and the germination of protein-containing seeds.§

In order to ascertain whether the enzyme be a tryptic one, a solution of it, or some of the more or less crude plant-extract, is added to a solution of peptone and placed in an incubator for some time, according to the strength of the solutions, kept at a temperature of 40°. A little toluol may be added as an antiseptic. To test, a few drops of the liquid are placed in a watch glass, acidified with acetic acid, and then a little chlorine water is added. The appearance of a marked yellow to pink coloration indicates the presence of tryptophane. If performed on a large scale, the liquid may be finally shaken up with amyl alcohol which dissolves the pink chlorine compound and eventually rises to the top. It may be separated by means of a small separating funnel and spectroscopically examined.

^{*} Dean: loc. cit. + Mendel: "Am. Journ. Med. Sci.," 1902.

[‡]Vines: "Ann. Bot.," 1905, 19, 171.

[§] Vines: id., 1902, 16, 1; 1903, 17, 237, 597.

An absorption band will be observed on the yellow side of the thallium line $(571 - 540\mu\mu)$.

Another test for tryptophane consists in mixing the suspected solution with a little glyoxylic acid and carefully adding concentrated sulphuric acid so that the latter forms a separate layer at the bottom of the test tube. After a short time a purple ring is produced at the junction of the two liquids, and on careful agitation the colour extends over the whole solution. If pepsin be used in the above experiments, it must be well washed in water and alcohol before use.

OUANTITATIVE DETERMINATION OF THE ACTIVITY OF PROTEASES.

Schultz* followed the course of the action of pepsin on egg albumen by precipitating out the albumen from time to time and examining the optical activity of the peptone solution.

Sjögvist, + on the other hand, measured the electrical conductivity of an albumen solution which was being hydrolysed by pepsin. †

Sörensen § found that he could obtain a measure of the amount of protein hydrolysed, by determining the number of free carboxyl groups in the mixture. By adding an excess of formalin, the free amino groups were neutralized; the carboxyl groups were then estimated by adding an excess of N/5 baryta solution and titrating back the excess by means of hydrochloric acid.

ZYMASE AND ALCOHOLIC FERMENTATION.

The formation of alcohol from fluids containing sugar has been known and practised for thousands of years, and the use of yeast in the manufacture of alcoholic beverages and of bread is an ancient industry. As is well known, when yeast is placed in a sugar solution, fermentation begins sooner or later, the principal end products being alcohol and carbon dioxide; substances other than ethyl alcohol, however, are formed,

^{*} Schultz: "Zeit, phys. Chem.," 1885, 9, 577. †Sjöqvist: "Skand. Arch. f. Physiol.," 1895, 5, 317. ‡Cf. Bayliss: "Arch. Sciences Biol.," St. Petersburg, 1904, 11 (supplem.).

[§] Sörensen: "Biochem. Zeit.," 1908, 7, 45.

especially glycerol, succinic acid and amyl alcohol,* the last more particularly in the fermentation of the sugars obtained from wheat and potato starch. Alcoholic fermentation is due to the activity of the enzyme zymase, which was first separated from the yeast cell by Buchner,† whose work ‡ marks the beginning of an epoch of vigorous investigations into this and kindred subjects.

It must, however, be remembered that many other enzymes besides zymase exist in *Saccharomyces*; e.g., diastase, invertase, trypsin and emulsin.

For laboratory purposes the fermentative activity of *Saccharomyces* may be quickly and conveniently illustrated by the use of Pasteur's solution, the composition of which is as follows:—

Ammonium tartrate, 50 grams Potassium phosphate, 10 grams Calcium phosphate, 1 gram Magnesium sulphate, 1 gram

These salts are thoroughly ground and mixed in a mortar, and I gram of the mixture together with 12 grams of glucose are dissolved in 70 c.c. of water, the yeast being added to the solution.

If cane sugar be used, marked fermentation will only begin after an interval of time has elapsed, during which the

*Amyl alcohol, using the term in its general acceptance, is a mixture of two isomeric primary alcohols, isobutyl carbinol ${\rm CH_3 \atop CH_3}$ CH . CH $_2$. CH $_2$ OH and CH.

secondary butyl carbinol CH₃—CH₂—CH—CH₂OH. The two substances together form "fusel oil," which is the harmful constituent of cheap spirit made from potatoes.

They appear to be produced from leucine (CH₃)₂CH—CH₂—CHNH₂COOH, CH₃

and isoleucine CH₃CH₂—CH—CHNH₂COOH, which are constituents of the protein molecule, by loss of CO₂ and replacement of the NH₂ group by OH (see p. 339). The mixture is optically active owing to the asymmetric carbon atom of the secondary butyl carbinol.

+For an account of our knowledge of alcoholic fermentation prior to 1897, see Green: "Nature," 21 April, 1898.

‡Buchner: "Ber. deut. chem. Gesells.," 1897, 30, 117, 1110; 1898, 31, 568; Buchner and Rapp: id., 1897, 30, 2668; 1898, 31, 209, 1084, 1090; 1899, 32, 127.

invertase secreted by the yeast converts the sucrose, which is not directly fermentable by yeast, into invert sugar.

Yeast, or yeast-juice, can set up fermentation in other substances besides glucose, such, for example, as galactose,* mannose, fructose,† sodium lactate,‡ and, according to Neuberg and Tir, common plant acids, fatty acids, glycerol and lecithin.

This fermentation, however, may not take place immediately on the introduction of the yeast to the particular substance: for instance, before yeast can ferment galactose, it must be educated with regard to this material by being cultivated for some time in a solution containing it. A yeast so educated yields a juice which can ferment galactose, the fermenting mixture, according to Harden and Norris, reacting with phosphate in a manner exactly similar to yeast-extract and glucose (see below); further, the process is accelerated by the addition of a small quantity of sodium arsenite.

In addition to ordinary alcoholic fermentation, yeast also exhibits the power of auto-fermentation. This is brought about at the expense of the reserve food-materials of the plant, chiefly glycogen, two enzymes being concerned in the process. Glycogenase changes the glycogen into sugar, which is then converted by zymase into alcohol and carbon dioxide, the rate of fermentation being dependent on the rate of sugar production by the glycogenase. Harden and Paine also found that the rate of auto-fermentation is greatly increased by the removal of water from the cell, which means, of course, a concentration of the cell sap. This may be accomplished by partial desiccation or by the use of dissolved substances which plasmolyse the cells. Alcohol in solutions above 10 per cent also have the same effect; on the other hand, salts which do not produce plasmolysis, even in concentrated solutions, such as urea, have no such accelerating effect.

^{*} Harden and Norris; "Proc. Roy. Soc., Lond.," B., 1910, 82, 645. + Harden and Young: id., 1909, 81, 336. ‡Kohl: "Beih. Bot. Centribl.," 1910, 29, 115. § Neuberg and Tir: "Biochem. Zeitschr.," 1911, 32, 323.

[#] Harden and Paine: " Proc. Roy. Soc., Lond.," B., 1912, 84, 448.

THE ISOLATION OF ZYMASE.

The following is the method pursued by Buchner in isolating zymase from Saccharomyces. One kilogram of compressed yeast is mixed with 250 grams of the infusorial earth known as kieselguhr and a quantity of fine quartz sand. mixture is ground in a mortar until the microscope shows the majority of the yeast cells to be broken. To this paste-like mixture are added 100 c.c. of water which is very thoroughly stirred in: the mass is then wrapped in a cloth, placed in a press and gradually subjected to a very high pressure— Buchner used a pressure as high as 500 atmospheres—the liquid extracted being collected in a glass vessel. The residue is then removed from the press, broken up, and again mixed with 100 c.c. of water and subjected to pressure. The extracts are united, shaken up with a little kieselguhr and filtered. The filtrate contains the zymase, but in an impure condition; it may be purified by precipitating with alcohol and dissolving the precipitate in water. The aqueous solution will not keep any great length of time, a character which is shared with most other enzymes when in aqueous solution; this phenomenon is termed hysteresis. It may, however, be preserved for a longer time—but not indefinitely—by drying the extract under reduced pressure, the solid substance so obtained being kept in a cold desiccator and dissolved in water as occasion demands.

In preparing extracts of yeast, it must be remembered that the potency of the extracts depends upon the physiological state of the yeast used. Thus, if brewers' yeast be taken from the wort whilst fermentation is at its height, a high quality zymase will be obtained; if, however, fermentation of the wort be over, the yeast taken from it will yield an extract of little or no fermenting power.

GENERAL CONSIDERATIONS.

The alcoholic fermentation of sugar by yeast may be represented by the equation:—

$$\mathrm{C_6H_{12}O_6} = 2\mathrm{CO_2} + 2\mathrm{C_2H_6O}$$

Recent investigations, however, have shown that the phenomenon is not quite so simple as this general statement indicates.

The names of Harden and Young particularly are associated with the problem, and a brief account of their work may be given.*

The alcoholic fermentation of glucose by yeast-extract is greatly increased by the addition to the fermenting liquor of yeast-juice, which has been boiled—so that all enzymes are destroyed—and filtered. After such addition, there is to begin with a greater evolution of carbon dioxide, which gradually diminishes to a rate which remains practically constant for several hours and usually is about equal to that given by an equal volume of the same yeast-extract and glucose to which boiled and filtered extract has not been added; but the diminution in the fermentation rate is slower than in the control, so that fermentation continues for a longer period, the extra amount of carbon dioxide evolved being directly proportional to the volume of boiled juice added. The accelerating factor is a phosphate, and analysis showed that the extra quantity of carbon dioxide evolved in the initial period of high evolution, when a phosphate or boiled extract is added, corresponds to the evolution of one molecular proportion of carbon dioxide for each atom of phosphorus added in the shape of It should thus be possible to separate yeast-juice into two complementary parts, either one of which depends upon the presence of the other in order that fermentation may take place. This was accomplished by filtering the expressed yeast-juice through a Martin gelatine filter under a pressure of 50 atmospheres. The residue remaining in the candle is inactive, + and so also is the filtrate which contains the active principle, which is neither destroyed by heat nor precipitated by ammoniacal magnesia mixture. This activating substance, or co-ferment, is a salt of hexose phosphoric acid.

The alcoholic fermentation of glucose therefore takes place

^{*}Harden and Young: "Proc. Roy. Soc., Lond.," B., 1906, 77, 405; 1906, 73, 369; 1908, 80, 299; 1909, 81, 336; 1910, 82, 321; 1911, 83, 451; "Centribl. Bakt.," 1910, 26, 178. Young: "Proc. Roy. Soc., Lond.," B., 1909, 81, 523; "Biochem. Zeit.," 1911, 32, 177.

⁺ The aqueous solution of the residue does not keep for long; it may, however, be preserved for some time by spreading it out on a clock glass and placing in a sulphuric acid desiccator. It dries to a brown, brittle substance which can easily be ground to a powder. In order to purify, the powder may be dissolved in water and once more filtered and dried as before.

in stages, the first of which is the formation of hexose phosphate which takes place during the first period of temporary acceleration

(1) ${}_{2}C_{6}H_{12}O_{6}+{}_{2}R_{2}HPO_{4}={}_{2}CO_{2}+{}_{2}C_{2}H_{6}O+C_{6}H_{10}O_{4}(PO_{4}R_{2})_{2}+{}_{2}H_{2}O$ This reaction only takes place provided that the enzyme and the co-ferment are present; soluble phosphates alone are unable to promote the fermentation in a mixture of the enzyme and glucose.

The hexose phosphate is continually being hydrolysed by an enzyme, hexose-phosphatase, yielding a free phosphate which again enters into combination with hexose:—

$$C_6H_{10}O_4(PO_4R_2)_2 + 2H_2O = C_6H_{12}O_6 + 2R_2HPO_4$$

The rate at which this second reaction takes place is the determinating factor in the fermentation rate when glucose is fermented by yeast-extract. There is an optimum concentration of phosphate which produces a maximum initial rate of fermentation; beyond this optimum a further addition of phosphate depresses the fermentative activity. If the available amount of phosphate in a mixture of sugar, ferment and coferment be very small, the total fermentation is greatly reduced, but if to such a mixture a little phosphate be added, there is an enormous increase, as much as 700 per cent, in the total fermentation, even after discounting an amount of carbon dioxide equivalent to the phosphate added.

With regard to other sugars, Harden and Young found that mannose and fructose are freely fermented by yeast-extract, fructose being fermented more quickly than mannose and mannose rather more quickly than glucose. Also the total weight of carbon dioxide given off from an excess of sugar by the action of a given volume of yeast-juice was slightly greater with fructose than with glucose, whilst that evolved from mannose was less than from glucose. No matter what sugar is used, glucose, fructose or mannose, the hexose phosphate is the same. The behaviour of fructose is qualitatively the same as glucose, but quantitatively there is a considerable difference. Thus the optimum concentration of phosphate for the fermentation of fructose is from 1.5 to 10 times as great as the optimum for glucose, and the maximum rate of fermentation of fructose is 2 to 6 times as great as that of glucose.

Fructose also behaves in the presence of other sugars as

an accelerating factor, for if the rate of fermentation of glucose or of mannose by yeast-extract is greatly lowered by the presence of a large excess of phosphate, the addition of a relatively small quantity of fructose brings about a marked acceleration in the fermentation. This is not due solely to the fermentation of the added fructose, for the amount of carbon dioxide evolved is much too great. This appears to be a specific property of fructose, for the phenomenon does not obtain when glucose is added to mannose or fructose, or by mannose when added to glucose or fructose under the proper conditions of concentration of phosphate in each case.

Commenting on this, Harden and Young observe that "this remarkable property of fructose, taken in connexion with the facts that this sugar in the presence of phosphate is much more rapidly fermented than glucose or mannose, and that the optimum concentration of phosphate for fructose is much higher than for glucose or mannose, appears to indicate that fructose when added to yeast-juice does not merely act as a substance to be fermented, but, in addition, bears some specific relation to the fermenting complex".

It is supposed that fructose forms a permanent part of the fermenting complex, so that a greater concentration of this sugar in yeast-extract leads to the formation of an increased quantity of complex. Thus, owing to the increased concentration of this active catalyst, the yeast-juice could bring about the reaction with sugar in the presence of phosphate at a higher rate and, at the same time, the optimum concentration of phosphate would become greater.

Harden and Young also find that the addition of a suitable amount of arsenate to a fermenting mixture of yeast-extract and sugar (glucose, fructose or mannose) causes a marked acceleration in the rate of production of alcohol and carbon dioxide, which is continued long after a chemical equivalent of carbon dioxide has been evolved. In this, the action of arsenate differs from that of phosphate and, further, the arsenate occurs in the free state throughout the period of fermentation. This increased rate of fermentation is due to the accelerating influence of the arsenate on the hexose-phosphatase; the arsenate, however, cannot replace phosphate in the fundamental reactions of alcoholic fermentation,

That phosphate is a necessity for alcoholic fermentation by zymase is generally agreed, but views other than the above have been put forward regarding the part played by it in fermentation.

Iwanoff,* for instance, considers that the phosphate formed is a triose phosphate, the formation of which is not necessarily accompanied by the evolution of carbon dioxide and alcohol, since the combination will take place when a phosphate is added to the filtrate of a solution of sugar which has been fermented by yeast-extract. He also found that the sugar obtained from the sugar phosphate is not fermented by living yeast. Iwanoff concludes that there are three stages in alcoholic fermentation: the sugar is first broken down into simpler sugars, then by the action of an enzyme, termed synthease, a triose phosphate is organized, which is then acted upon by alcoholase to form carbon dioxide, etc.

These views are not agreed with by Harden and Young,† who criticize the methods employed by Iwanoff.

According to Buchner, lactic acid is an intermediate product of fermentation; in the first place the glucose under the influence of zymase is converted into lactic acid, which is then attacked by another enzyme, the action giving origin to carbon dioxide and alcohol.

Kohl,‡ however, points out that lactic acid is not fermented by zymase, by compressed yeast nor by bottom yeast; indeed I per cent lactic acid is sufficient to stop the auto-fermentation of yeast and to reduce greatly the fermentation of glucose. On the other hand, zymase will ferment sodium lactate, which indicates that if lactic acid is an intermediate product of fermentation, according to Buchner's view, a salt rather than the acid must be formed.

In yeast-extract Kohl found an enzyme, catalase, which was capable of oxidizing phenols. The yeast-extract on filtering produces lactic acid in the presence of glucose, and the acid in the presence of zymase is converted into alcohol and carbon dioxide; if, however, zymase be not present, oxidation may go further and other acids be produced.

^{*} Iwanoff: "Centrlbl. Bakt.," 1909, 24, 1.

⁺Harden and Young: "Centrlbl. Bakt.," 1910, 26, 178.

[‡] Kohl: "Beili, bot, Centrlbl.," 1910, 20, 115.

Briefly put, he considers that the glucose, by the action of catalase, is converted into lactic acid which is operated upon by zymase, so that alcohol and carbon dioxide are produced.

Other opinions have been put forward, but as the evidence is not always sufficiently strong to stand the weight of the theory, it is not proposed to consider them here.

Zymase-like enzymes are not restricted to the yeasts; such bodies have been identified in other Fungi, such as *Mucor stolonifera** and *Aspergillus niger.*†

IDENTIFICATION OF ETHYL ALCOHOL.

The following tests may be employed for the identification of ethyl alcohol in fermented liquors, etc.:—

- 1. The presence of ethyl alcohol, except in very dilute solutions, is usually betrayed by the smell.
- 2. To the suspected solution add an equal volume of a solution of iodine in potassium iodide, and then carefully add just sufficient caustic potash to decolorize the iodine; on gently warming the solution, a smell of iodoform is produced and a yellow crystalline precipitate is formed if alcohol be present. If alcohol is present in abundance, it may keep the iodoform in solution, and so prevent the formation of a precipitate; this may be remedied by adding water.

It is to be observed that this reaction is not given by methyl alcohol, but is given by a number of other substances, such as acetic aldehyde and acetone. Ethyl alcohol will, however, not produce iodoform in the presence of ammonia, whereas acetone will.

- 3. Solutions of alcohol mixed with an equal volume of concentrated sulphuric acid, and warmed with a few crystals of sodium acetate, evolve a pleasant fruity odour of ethyl acetate.
- 4. Warmed with potassium dichromate solution and a little dilute sulphuric acid, alcohol is oxidized to acetaldehyde, which may be recognized by its smell.

^{*} Kostytschew: "Ber. deut. bot. Gesells ," 1904, 22, 207. † Maximow: id., 1904, 22, 225.

QUANTITATIVE DETERMINATION OF THE ACTIVITY OF ZYMASE.

The amount of carbon dioxide evolved, either by volume or by weight, may be taken as a measure of the enzymic activity.

For the estimation of the volume and the rate of evolution of gas given off during the fermentation of sugar by yeast-juice, Harden, Thompson and Young* give the following method (Fig. 5) which yields satisfactory results, provided

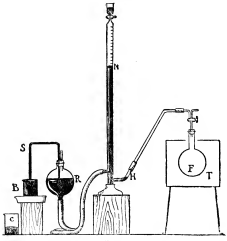


FIG. 5.

that not more than 30 c.c. and not less than '5 c.c. of gas is evolved in five minutes.

The fermenting mixture is placed in the flask F, of 100 to 200 c.c. capacity, which is kept at a constant temperature by means of a thermostat, T. The flask is fitted with a three-way tap, which is connected by means of rubber and glass tubing to a Schiff nitrometer, N, of 100 c.c. capacity and graduated to 2 c.c. The nitrometer is connected with a reservoir, R, which is fitted with a bung, through which passes

^{*} Harden, Thompson and Young: "Biochem. Journ.," 1911, 5, 230.

a siphon tube, S; the other end of this tube dips into a beaker, B, which stands in a dish provided with a lip, so that the excess of mercury which falls from B is ultimately discharged into a second beaker, C. Any mercury displaced from N collects into the reservoir R, and is siphoned off into D, from whence it overflows into the shallow dish, and is finally collected in C. The mercury at R and H, therefore, remains level with the mercury in B. This adjustment is made once for all, so that the mercury extends to the bend of the tube H, and thus the pressure in the flask is maintained at atmospheric pressure.

The volume of the gas given off is read on N at the reduced pressure corresponding to the height of the column of mercury NH, and is corrected by means of a table which is drawn up as follows. The height of each graduation mark of the nitrometer above H is measured, and subtracted from 760 mm., the normal pressure of the atmosphere; the corrected volume corresponding with each graduation is then calculated. When great accuracy is required the reading must be corrected for the atmospheric pressure obtaining at the time of the experiment.

In starting an experiment the nitrometer is filled with mercury, and the siphon adjusted in R and started by means of a pressure bulb which is attached to the short tube passing through the stopper of the reservoir. The fermenting mixture is placed in the flask F, together with a little toluene as an antiseptic. When the temperature has reached that of the thermostat and is constant, the flask is removed, filled with carbon dioxide and thoroughly shaken, the process being repeated until the liquid is saturated. Owing to the fact that the liquid easily becomes supersaturated the flask must be well shaken for half a minute before a reading is taken. The flask is replaced in the thermostat for a minute, in order to raise the temperature to the proper degree, and then the volume of gas collected in the nitrometer is read.

At the expiration of the desired interval of time from the last shaking the flask is again thoroughly shaken, after which the volume of gas evolved during the interval is read.

When the nitrometer is filled with gas, the tap of F is closed, the siphon removed, the reservoir filled with mercury

from C and the gas displaced by raising the reservoir. To recommence the collection of gas, the siphon is replaced and started with a pressure bulb, and then the tap of F is opened.

By means of the mercury cup on the top of the nitrometer, samples of the gas may be removed for analysis.

It is to be noted that as rubber tubing is permeable to carbon dioxide as little as possible should be used for the connexions between F and N.

OCCURRENCE OF ALCOHOLS IN PLANTS.

Methyl Alcohol has been found to occur in the aqueous distillates and in the essential oils of a very large number of different plants, amongst which might be mentioned Juniperus Sabina, Zea Mais, Lolium perenne, Iris germanica, Euonymus europaea, Thea sinensis, Eugenia caryophyllata, Carum carvi, Anthriscus cerefolium, etc.

Ethyl Alcohol is not quite so widely distributed as methyl alcohol, but occurs in distillates from Cananga odorata (Ylang Ylang), Pyrus Malus, Mespilus germanica, Eucalyptus, Anthriscus cerefolium, Pastinaca sativa, Vaccinium Myrtillus, Betula alba. etc.

Mention also should be made of the occurrence of this alcohol, together with lactic acid and acetone * in some cases, in the higher plants especially during anaerobic respiration. Stoklasa,† for instance, found that this substance together with acetic and formic acids was produced during anærobic respiration of potatoes and seeds. Indeed, many consider that alcoholic fermentation is the first expression of respiration, and whether alcohol is formed or not depends upon the conditions; thus under normal conditions in the presence of oxygen the first products are oxidized before the alcohol stage in the process is reached, or the alcohol may be used up in anabolic processes as soon as it is formed, or it may be oxidized to water and carbon dioxide—the normal end products of ærobic respiration.‡

^{*} Palladin and Kostytschew: "Ber. deut. bot. Gesells.," 1906, 24, 273.

⁺ Stoklasa: id., 1904, 22, 358; "Centr. f. Bakter. u. Parasit.," 1905, II, 31, 86. Godlewski and Polzeniusz: "Bull. Acad. Sci., Cracow," 1901, 227; Stoklasa, Jelinek and Vitek: "Beitr. z. chem. Phys. u. Path.," 1903, 3, 460,

[‡]See Kostytschew: "Ber. deut. bot. Gesells.," 1908, 26, 565.

Amyl Alcohol has been identified in the essential oils of geranium, eucalyptus, lavender, peppermint and chamomile.

Several unsaturated alcohols, such as citronellol $C_{10}H_{20}O$, geraniol and linalool, both of the formula $C_{10}H_{18}O$, occur in essential oils, such as rose oil and oil of bergamot, while amongst the alcohols belonging to the aromatic series must be mentioned cinnamic alcohol, benzyl alcohol, menthol, borneol, etc. Other monohydric alcohols, with the exception of phytosterol (see p. 17), are of comparatively rare occurrence.

Examples of polyhydric alcohols occurring in plants are mannitol, sorbitol, and dulcitol, isomeric substances of the formula—

СН2ОН СНОН СНОН СНОН СН2ОН

Mannitol occurs to the extent of about 40-50 per cent in manna, the dried sap of Fraxinus Ornus, and also in celery, Syriaga vulgaris, etc. etc. Sorbitol occurs in the berries of the mountain ash, Pyrus Aucuparia. Dulcitol occurs in the cortex of Euonymus europaea and in the bark of Euonymus atropurpurea.

Adonite,* C₅H₁₂O₅ or CH₂OH CHOH CHOH CHOH CH₂OH,

is a pentahydric alcohol occurring in *Adonis vernalis*. According to Treboux, † it is converted by the plant into starch. Adonite has a sweet taste, and is used in bacteriological media.

Of recent years a number of dihydric alcohols of high molecular weight have been found to occur in plants. They belong to different series whose general formulæ are—

$$C_nH_{2n-6}O_4,\,C_nH_{2n-8}O_4,$$
 and $C_nH_{2n-10}O_4.$

Trifolianol, $C_{21}H_{34}O_2(OH)_2$, isolated by Power and Salway,‡ from red clover leaves, may be taken as an example of the first group, while *Bryonol*, $C_{22}H_{34}O_2(OH)_2$, obtained by Power and Moore § from Bryony root, and *Calabarol*, $C_{22}H_{34}O_2(OH)_2$, isolated by Salway|| from Calabar beans, are representatives of the second and third groups respectively.

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* Merk: "Chem. Zentr.," 1893, 344.

† Treboux: "Ber. deut. bot. Gesells.," 1909, 27, 428.

‡ Power and Salway: "J. Chem. Soc., Lond.," 1910, 97, 249.
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[§] Power and Moore: "J. Chem. Soc., Lond.," 1910, 97, 249.

^{||} Salway: "J. Chem. Soc., Lond.," 1911, 99, 2155.

Of the polyhydric alcohols Inosite is of particular interest, and may, therefore, receive more detailed consideration.

INOSITE.

Inosite, which has the formula $C_6H_{12}O_6$, is isomeric with the hexoses, and, like these substances, has a sweet taste; for these reasons it was at one time thought to be a true sugar and was called muscle sugar owing to its occurring in muscle.

Inosite is, however, not a carbohydrate at all but a polyhydric alcohol derived from benzene and having the constitution—



Besides being found in muscle, inosite is of common occurrence in plants, in the leaves, especially when young, of *Vitis*, *Juglans*, etc.; in the roots and rhizomes of very many plants; in various seeds and fruits, e.g. *Phaseolus*, *Pisum*, and other leguminous seeds, *Vitis*, various cereals, and oily seeds, such as mustard.

It may be looked upon as a plastic substance since Maquenne has found that it disappears from the young fruits of *Phaseolus* as ripening proceeds.

Preparation.

The method of separation of inosite from the plant juices is based on the fact that it forms a compound with lead oxide.

The sap is expressed from the organ, or, if this be impracticable, the parts are ground up very thoroughly with water. The liquid is then filtered and, if it gives an acid reaction, is neutralized by the addition of baryta water.

A solution of basic lead acetate is then added until no more precipitate comes down. The precipitate consists of a compound of inosite with lead oxide (2C₆H₁₂O₆ 5PbO), and is filtered off, then washed and suspended in water, and saturated with a current of sulphuretted hydrogen. The lead sulphide

is filtered off and the filtrate evaporated on a water bath to the consistency of a syrup. On the addition of alcohol, containing one-tenth of its volume of ether, inosite is deposited in prismatic crystals.

Inosite has a sweet taste, is soluble in water but insoluble in alcohol and ether. It crystallizes in prisms, and does not give any reactions characteristic of true hexoses. For instance, it is not fermentable, it does not reduce Fehling's solution, and its solution does not give a brown coloration with potash.

Identification.

- 1. When moistened with a little dilute nitric acid, then evaporated almost to dryness, and made alkaline with ammonia, the addition of a few drops of chloride of calcium produces a rose-red coloration.
- 2. A solution of inosite evaporated to dryness with a few drops of mercuric nitrate produces a yellow stain which on heating turns red.
 - 3. Solutions of inosite are not optically active.

With regard to its significance in the plant there is evidence to show that inosite is a transitory substance and is used up in the synthesis of other substances.

According to Posternack * a large amount, 80-90 per cent, of the phosphorus of certain seeds exists in the form of phytin; it occurs, for instance, in the globoid portion of aleurone grains, and the seeds which contain it also possess an appropriate enzyme phytase for its decomposition into phosphoric acid and inosite.†

With regard to the formation of phytin little is known; Posternack considers that it is formed by the combination of formaldehyde, produced in the early stages of photosynthesis with phosphoric acid.

The tenability of this opinion is obviously bound up with the formation of formaldehyde in green leaves (q.v.).

Phytin appears to be an acid calcium and magnesium salt

^{*} Posternack: "Compt. rend.," 1903, 137, 202, 337, 439. † Cf. Suzuki, Yoshimura and Takaishi: "Bull. Coll. Agric., Tokyo," 1907, 7, 503. See also Rose: "Biochem. Bull.," 1912, 1, 428.

of inosite phosphoric acid which is a condensation compound of inosite with six molecules of phosphoric acid.*

MANUFACTURE OF ETHYL ALCOHOL.

The action of yeast on sugar is made use of in the manufacture of ethyl alcohol, which substance is prepared from potatoes, rice, and other grains rich in starch. The manufacture from potatoes is carried out as follows: Potatoes are heated in closed vessels to 125-135° by means of superheated steam under a pressure of about 3 atmospheres; by suddenly releasing the pressure the potatoes are burst, and are thus obtained in a finely divided state. The whole mass is then thoroughly stirred up with malt at a temperature of about 60°, whereby the starch undergoes hydrolysis with formation of maltose and dextrin.

$$(C_6H_{10}O_5)_n + H_2O \rightarrow C_{12}H_{22}O_{11} + (C_6H_{10}O_5)_x$$

Starch Maltose Dextrin

After about one and a half hours the mixture is rapidly cooled to 15° and mixed with yeast; fermentation at once sets in, accompanied by a considerable evolution of heat; the mixture is therefore cooled artificially, so that the temperature is maintained steady at about 27°:5-30°.

During this time the maltose is converted first into dextrose and then into alcohol and carbon dioxide according to the equations:—

$$\begin{array}{c} C_{12}H_{22}O_{11}\,+\,H_2O\,=\,2C_6H_{12}O_6\\ C_6H_{12}O_6\,=\,2C_2H_5OH\,+\,2CO_2 \end{array}$$

In order to convert the dextrin, which would otherwise be lost, into a fermentable substance, the temperature towards the end is maintained at about 26-29° in order to give the malt a further opportunity of hydrolysing the dextrin to glucose, and so rendering it capable of being fermented by yeast. When the fermentation is completed after about three days, the mixture contains about 13 per cent of alcohol by volume; by distilling the mixture through a fractionating

^{*} Cf. Neuberg: "Biochem. Zeitschr.," 1908, 9, 557; Winterstein: "Zeitschr. physiol. Chem.," 1908, 50, 118. See also Plimmer: "Biochem. Journ.," 1913, 7, 43.

column, so much of the water is removed that the distillate contains about 80 to 95 per cent of alcohol.*

No amount of fractional distillation without dehydrating agents will produce alcohol containing less than 4:43 per cent by weight of water, since such alcohol gives a constant boiling mixture.

Alcohol containing 0.5 per cent or less of water is, in commerce, known as absolute alcohol, although in a scientific laboratory the term is only correctly applied to alcohol which is quite free from moisture; such alcohol can only be obtained by careful fractionation from freshly burnt quicklime.† If the alcohol is dehydrated over quicklime to which a little barium oxide has been added, complete dehydration is marked by the formation of a yellow colour due to the production of barium ethylate, which can only be formed in the absence of any trace of moisture.

A delicate test for the detection of traces of moisture in alcohol consists in adding a few drops of the sample to a solution of liquid paraffin in anhydrous chloroform; if there is any moisture present, a turbidity will be at once produced.

OXIDASES.

The oxidases are enzymes which have the power of oxidizing various aromatic compounds and chromogens, which action is frequently indicated by a change in colour. This change in colour in vegetable tissues on exposure to air is an everyday phenomenon; the exposed surfaces of a bitten apple, especially cider varieties, will rapidly turn brown; similarly the fruit-body of *Boletus* quickly assumes a prussian-blue colour on being broken. The darkening in the colour of raw rubber is also due to an oxidase which is associated with the protein of the coagulated latex.‡

These changes are often of considerable economic importance; thus the discoloration of sap wood markedly depreciates

^{*} The residue remaining after distillation contains, in addition to the solid unfermentable materials, a certain amount of other soluble products of fermentation, such as glycerol and succinic acid; it is used as a cattle food.

[†]Occasionally the last traces of moisture are removed by treating the alcohol with sodium wire.

[#] Spence: "Biochem. Journ.," 1908, 3, 165, 351.

the value of the timber,* while the lacquer industry of China and Japan has been built up on the facts relating to the action of the oxidase, laccase, on the expressed sap of species of *Rhus.* (See below.)

Oxidases are very widely distributed in the vegetable kingdom; in the higher plants they may occur in any organ—stem, root, leaf, laticiferous tissue, petals, and fruits.

Several oxidases have been distinguished, e.g. laccase, which has already been mentioned; tyrosinase, which oxidizes tyrosine into homogentisinic acid (p. 359); olease, from olives, which can oxidize fats into simpler fatty acids, † and others which oxidize sugars into carbon dioxide and water.‡

The action of oxidases may be illustrated by a brief reference to laccase, an enzyme which was first investigated by Yoshida.§ The latex of many species of *Rhus* rapidly turns brown and finally black on exposure to the atmosphere; if the juice be evenly spread out, the final product is black and shiny. The extract of the plant contains urushic acid (laccol) which is oxidized into oxyurushic acid:—

$$\mathbf{C_{14}H_{18}O_2 + O} = \mathbf{C_{14}H_{18}O_3}$$

The action takes place best at 20° C. in the presence of moisture and oxygen; at higher temperatures it is destroyed, at 63° according to Yoshida, and at 70° according to Bertrand. Bertrand \(\) also has given much attention to this oxidase, and the most important fact ascertained by him in this connexion is that the presence of manganese is all-important. He found that the activity of the ferment is directly proportional to the amount of the metal present, which acts as a co-enzyme (p. 355). But whether manganese is essential for all oxidase reactions is uncertain, for Bach \(\) states that he has prepared a tyrosinase which will oxidize tyrosine in the absence of manganese and of iron.

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* Bailey: "Bot. Gaz.," 1910, 50, 142.
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⁺ Tolomei: "Chem. Centrbl.," 1896, 1, 879.

[‡]See Palladin: "Zeit. physiol. Chem.," 1906, 47, 407. §Yoshida: "J. Chem. Soc., Lond.," 1883, 43, 472.

^{||} Bertrand: "Compt. rend.," 1895, 120, 266; 1895, 121, 166; 1896, 122, 1132; 1896, 123, 463; 1897, 124, 1032, 1355.

[&]quot; Bach: "Ber. deut. chem. Gesells.," 1910, 43, 362.

ISOLATION OF OXIDASES.

The isolation of oxidase may be a difficult matter when it exists in a tissue together with its substrate and other enzymes. Bourquelot and Bertrand give the following method for Fungi such as Russula. The tissue is chopped up, extracted with water—which may be warmed—and filtered as quickly as may be. The filtrate is then poured into an excess of strong alcohol, whereby the enzyme is precipitated. The precipitate is then filtered off and dissolved in water.

When the oxidase is extracted together with other oxidizing enzymes, separation may be effected, at any rate in part, by adding to the aqueous solution of the oxidase, or to the aqueous extract of the plant, two volumes of absolute alcohol for each volume of extract. The oxidase will be precipitated, whilst the other enzymes will remain in solution.*

PEROXIDASE.

Peroxidases which split oxygen from hydrogen peroxide, organic peroxides, potassium permanganate, etc., are often associated with other oxidases, and are very widely distributed; indeed, they have even been described as occurring in coal.†

Preparation of Peroxiduse.

Appleman ‡ recommends the following method of preparing peroxidase. Potato tubers are grated into a pulp, which is thoroughly mixed with calcium carbonate in order to neutralize any acids. The mass is then ground with quartz sand in a mortar for about two minutes and filtered through butter muslin. The extract contains the peroxidase together with oxidase; the latter may be removed by raising the temperature to 70° for ten minutes, whereby the oxidases are coagulated, or, according to Gruess, any oxidase present may be destroyed by adding acetone which does not affect the peroxidase. In many experiments where only a dilute solution of peroxidase is required—I c.c. of extract to 300 c.c. of water

^{*} Aso: "Bull. Coll. Agric. Imp. Univ., Tokyo," 1902, 5, 207. + Stoklasa, Ernst and Chocensky: "Ber. deut. bot. Gesells.," 1907, 25, 38.

[‡] Appleman: "Bot. Gaz.," 1911, 52, 306. § Gruess: "Ber. deut. bot. Gesells.," 1903, 21, 356.

—the heating may be dispensed with, as the amount of oxidase is so very small.

The method followed by Gruess* is somewhat different. The potatoes are sliced into absolute alcohol, and the oxidases destroyed by heating to 70° for ten minutes. The slices are allowed to remain twenty-four hours in absolute alcohol, which should be changed at least three times. The material is then superficially dried with filter paper and covered with ether for a few minutes. The dehydrated slices are then freed from the alcohol and ether by placing in a vacuum desiccator, after which they may be ground up in a mortar. Before use, I gram of the powder is ground with sand and 25 c.c. of water, and then filtered.

This method is criticized by Appleman, who points out that in the process of drying, the activity of the peroxidase is greatly impaired, and also that the presence of coagulable proteins interferes with the stability of the peroxidase activity, besides causing a low yield of enzyme.

Identification.

- 1. Guaiacum tincture in the presence of oxidase turns blue provided oxygen be present.
- 2. In cases where the blueing of the guaiacum tincture does not take place immediately, the addition of hydrogen peroxide may bring it about.
- 3. Tetramethyl-p-phenylenediamine in the presence of hydrogen peroxide gives a deep violet colour with an oxidase.
- 4. Peroxidases set free oxygen from hydrogen peroxide and other peroxides.

For comparative experiments with peroxidase, Gruess uses I gram of pulverized powder, prepared as above, which is mixed and ground with sand with 25 c.c. of water. For the test, 5 c.c. of the filtrate is mixed with '5 c.c. of guaiaconic acid dissolved in alcohol, and 'I c.c. of a '5 per cent solution of hydrogen peroxide.

Appleman, for comparative tests, allows a definite quantity of the extract (see above) to act on a definite volume of guaiaconic acid solution in the presence of hydrogen per-

^{*}Gruess: "Zeit. Pflanzenkrank.," 1910, 25, 115.

oxide, the test tube being kept at a constant temperature whilst the experiment is going on. For the comparison, a standard blue aqueous solution of indigo carmine is made; the time required for the blue of the guaiacum mixture to match the colour of the standard blue is taken as the index of the peroxidasic activity.

It should be remarked that, according to Aso, the presence of certain substances, e.g., tannin or sodium fluoride, interferes with the colour reactions normally given by oxidases.

GENERAL CONSIDERATIONS.

Up to comparatively recent times an oxidase was considered to be a single enzyme, but according to Bach and Chodat,* what used to be termed oxidase is really a mixture of peroxidase and peroxide. According to them, there are three categories of oxidizing ferments.

- (a) Oxygenases which produce the peroxide.
- (b) Peroxidases which transfer oxygen from the peroxide to the substance to be oxidized.
- (c) Catalases which destroy peroxides so that oxygen is given off.

In the colour reactions mentioned above two actions are possible. Either the plant juice, e.g. of the potato, gives the blue coloration with the guaiacum tincture alone, or, the blue colour will not occur, as, for example, in the sap of the cucumber, unless a peroxide, such as hydrogen peroxide, be added.

On Bach and Chodat's hypothesis, there are present in the potato oxygenase, peroxidase and peroxide; the peroxidase transfers oxygen from the peroxide to the guaiacum, and the oxygenase re-oxidizes the reduced peroxide. This may be termed the direct action.† On the other hand, in the cucumber juice, only peroxidase is present, so that in order to obtain the blue reaction with guaiacum, hydrogen peroxide, or other peroxide, must be added. This is the indirect action.

This idea has been accepted by Palladin,‡ who considers

^{*}Bach and Chodat: "Biochem. Centrbl.," 1903, 1, 416; Bach: "Ber. deut. chem. Gesells.," 1906, 39, 2126; 1907, 40, 230; 1908, 41, 216.

[†] Wheldale: "Proc. Roy. Soc., Lond.," B., 1911, 84, 121.

[‡] Palladin: "Ber. deut. bot. Gesells.," 1906, 24, 97.

that normal respiration depends upon the presence of an oxidizable substance, oxygenase and peroxidase.

Peroxidases can practically always be found in living plant members, but the oxygenases are less stable and are quickly decomposed, giving origin to some of the respiratory carbon dioxide. The amount of these enzymes varies with the stage of development of the plant; thus in the embyro, oxygenase is at its minimum, but increases with the development of the plant and then diminishes as the growth of the organ ceases.

On the other hand, according to Porodko,* oxidases play scarcely any part in respiration.

The views of Bach and Chodat are not universally held; thus Moore and Whitley,† as a result of a number of experiments, have arrived at the conclusions that the sole difference between the various plant extracts, etc., which show an oxidizing action, consists in the presence of a small variable amount of peroxide which is chemically unstable. Juices possessed of such oxidizing properties have one type of ferment, a peroxidase, which acts only in the presence of peroxide, which, if not present in the natural extract, must be added. There is no proof of the existence of any other type of enzyme, such as oxygenase, engaged in oxidation processes. Thus the oxidases are brought into line with hydrolytic enzymes concerned in the phenomena of digestion, etc.:—

	Hydrolysis.	Oxidation.		
Substrate	Carbohydrates, fats, proteins	Oxidizable substances, e.g. phenols and chromogens		
Combining body ("combinate")	Water (finally)	Oxygen which is yielded by hydrogen peroxide or organic peroxides		
Catalyst	Diastase, zymase, etc.	Peroxidase, tyrosmase, etc.		

They further point out that any substance containing a peroxide linkage will activate a peroxidase just as any type

^{*} Porodko: "Beih. Bot. Centrbl.," 1904, 16, 1.

⁺ Moore and Whitley: "Biochem. Journ.," 1909, 4, 136.

of acid or alkali, which increases hydrogen or hydroxyl ion concentration, will activate a hydrolytic enzyme.

The reason why a plant extract containing oxidases will no longer give the guaiacum reaction when heated to 60° is that the peroxide, originally present in the juice, is destroyed, but not the peroxidase. So that although the heated juice is inactive, its oxidizing activity can be restored by the addition of a peroxide.

The work of Moore and Whitley is corroborated by Wheldale,* who finds that the power of the direct action, but not the indirect, is accompanied by the formation of a brownish pigment when the part is injured or subjected to the action of chloroform vapour. This action is common in the Compositæ, Umbelliferæ, Labiatæ, Boraginacæ, and certain genera of the Scrophulariacæ, Rosacæ, Leguminosæ, and Ranunculacææ. In the Cruciferæ, Caryophyllacææ, Crassulacææ and Ericaceæ the action is either absent or very rare. In such cases she finds the direct action to be due to pyrocatechin which, on exposure to air, rapidly oxidizes and then acts as an organic peroxide, thus enabling the peroxidase, which is almost universally present, to transfer oxygen to the oxidizable substance.

In addition to oxidative processes carried out in the plant by these enzymes, they may play an important part in the preparation of raw food-material in the soil. Thus Schreiner and Reid + as the result of their experimental work have come to the conclusion that enzymes, principally peroxidases, are excreted by the roots, and so oxidize organic substances in the soil. Oxidation was found to be most potent in the roothair regions and those places where the secondary roots are developing. The older parts of the root and also the root-cap do not produce these enzymes. The degree of the activity of the enzymes in the soil depends upon the character of the soil; thus the presence of sodium nitrate accelerates the oxidative processes, and, in brief, the most productive soils show the most vigorous oxidation. On the other hand, the reverse obtains in poor unproductive soils which contain substances which interfere with the process. These deleterious agents

^{*}Wheldale: "Proc. Roy. Soc., Lond.," B., 1911, 84, 121. +Schreiner and Reid: "Bot. Gaz.," 1909, 47, 355. See also Schreiner and Sullivan: id., 1911, 51, 156, 273.

may be removed sometimes by suitable treatment, e.g., the use of absorbing agents such as lamp-black.

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INDEX.

ABIURETIC derivatives of proteins, 313. | Aldol condensation, 150; of formalde-Abrus precatorius, 274. hyde, 151. Absolute alcohol, 390. Aleppo galls, 212. Acacia, 195, 211. Aleurone grains, 301. - catechu, 126, 217, 246. Algæ, 110, 115, 127, 128, 138, 157, 194. leucophlæa, 126. 229, 257, 258. Accelerators, 354. Algarobilla, 210, 216. Acer pseudoplatanus, 279. Alisma, 115. Acetal, 149. - plantago, 98. Acetaldehyde cyanohydrin, 149. Alkali albumin, 327. Alkaloidal reagents, 270. Acetone, occurrence in plant, 285; cyanhydrin, 172, 337. Alkaloids, microchemistry, 271; occur-Acetyl cellulose, 135. rence, 265; origin, 272; physiology, Acetyl value, determination, 33; of ger-280. minating sunflower, 37; table, 34. Allihn's tables for gravimetric estima-Achroodextrin, 107, 110. tion of glucose, go. Acid albumin, 327. Allium, 97, 115, 154; germination, 40. - amides, 315. Сера, 101. Aconitine, 269, 271. Allyl isothiocyanate, 188. Aconitum, 266. Almond, 36, 174; amount of fat, 2. - Napellus, 271. - oil, 19, 32. Acorus calamus, 275. Aloe, 252. Acrolein, 21, 44. Althaein, 249. Althaca rosca, 249. Activators, enzymic, 354. Adamkiewicz's reaction, 307. Amanita muscaria, 274, 275. Adenine, 278, 281. Amaryllidaceæ, 115. Adipocellulose, 132, 137. Amide nitrogen, 318. Adonis vernalis, 386. Amides, 315. Amino acids, 315; occurrence in plant, Adonite, 386. 321; synthesis in plant, 337, 338. Adsorption, 295. selective, 297. Ammonium, as source of nitrogen, 333; Aesculus Hippocastanum, 45, 197, 321. oxidation, 335. Ampelopsis hederacea, 255. Aetiophyllin, 232. Aetioporphyrin, 233. Amphoteric electrolytes, 311. Agavose, 53, 73. Amygdalin, 169, 171, 172, 181; hydrolysis, 182; preparation, 181. Amyl alcohol, 338, 339, 375, 386. Agaracineæ, 194. Agaricus muscarius, 49. Agave americana, 73. Amylase, v. Diastase. mexicana, 153. Amylo-cellulose, 96, 100, 101, 108. Agrostis, 120. Amylo-dextrin, 100, 107, 109. Amyloid, 123, 135. Alanine, 315. Albizzia, 126. Amyloin, 107, 108. Albumins, 322. Amylo-pectin, 100, 101, 108. Albumose, 327. Aniylose, 100, 101, 104, 108. Alcogel, 284. Amylum, see Starch. Alcohol, absolute, 390. Ananas sativa, 98, 370. Alcoholic fermentation, 374-85. Angiosperms, 194, 265, 274. Alcohols, occurrence, in plant, 385. Anthericum, 115. Alcosol, 284. Anthoceros, 127.

Anthocyanidin, 249. Anthocyanin, 244, 248, 249, 252; and flavones, 244; and sugars, 254; and tannin, 255; extraction, 251; physiology, 256; properties, 254; reactions, 249, 256. Anthocyanins and anthoxanthins, 251. Anthoxanthin, 244, 247, 248. Anthicease erefolium, 385. — vulgaris, 288. Anti-enzymes, 356. Aphis chinensis, 212. Apignenin, 246. Apocynaceæ, 205. Apomorphine, 269. Apple, 68, 121, 128. — seed oil, 20. Apposition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 331. Arabic acid, 124, 125. Arabinon-trigalactan-geddic acid, 726. Arathis, germination, 41. Arachis oil, 19, 22. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Arachiaceæ, 138. Arbutin, 172, 174, 121, 122, 276, 306, 324, 346, 369. Beetword, 68, 71, 74, 121, 122, 276, 306, 324, 346, 369. Bear, 132, 371. Beech-nut oil, 20, 32. Beeswax, 43. Beetword, 68, 71, 74, 121, 122, 276, 306, 324, 346, 369. Bean, 132, 371. Beech-nut oil, 20, 32. Beeswax, 43. Beetword, 68, 71, 74, 121, 122, 276, 306, 324, 346, 369. Bean, 132, 371. Beech-nut oil, 20, 32. Beeswax, 43. Beetword, 68, 71, 74, 121, 122, 276, 306, 324, 346, 369. Been 132, 346, 369. Beetword, 68, 71, 74, 121, 122, 276, 306, 324, 346, 369. Bear, 132, 371. Beech-nut oil, 20, 32. Beetword, 68, 71, 74, 121, 122, 276, 306, 324, 346, 369. Bear, 132, 371. Beech-nut oil, 20, 32. Beetword, 68, 71, 74, 121, 122, 276, 306, 324, 346, 369. Bear, 132, 371. Beech-nut oil, 20, 32. Beetword, 59; yield of sugar, 69. Beeswax, 43. Beetword, 67, 41, 128, 252, 279, 347. Beeswax, 43. Beetword, 67, 41, 128, 252, 279, 347. Beeswax, 43. Beetword, 67, 41, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetword, 35. Beetword, 68, 71, 724, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetword, 35. Beetword, 68, 71, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetword, 35. Beetword, 68, 71, 128, 252, 279, 347. Beetword, 68, 74, 128, 252, 279, 347. Beetword, 68, 74, 128, 252, 279, 347. Beetword, 68, 74, 128, 252,		
flavones, 244; and sugars, 254; and tannin, 255; extraction, 251; physiology, 256; properties, 254; reactions, 249, 256. Anthocyanins and anthoxanthins, 251. Anthoxanthin, 244, 247. Anthrisics exerfolium, 365. — wilgaris, 288. Antiseptics and enzymes, 366. Aphis chinensis, 212. Apigenin, 246. Apocynacese, 205, Apomorphine, 269. Apple, 68, 712, 712. Beassoric acid, 126. Bean, 132, 371. Bean, 132,	Anthocyanidin, 249.	
tannin, 255; extraction, 251; physion logy, 256; properties, 254; reactions, 249, 256. Anthocyanins and anthoxanthins, 251. Anthoxanthin, 244, 247. Anthriscus eerefolium, 385. — wulgaris, 288. Anti-enzymes, 358. Anti-enzymes, 358. Anti-enzymes, 358. Anti-enzymes, 358. Anti-enzymes, 358. Anti-enzymes, 358. Anti-enzymes, 366. Aphis chinensis, 212. Apigenin, 246. Apocynaceæ, 205. Appomorphine, 269. Apple, 68, 721, 128. — seed oil, 20. Apposition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, Arabin, 124, 125. Arabinon-trigalactan geddic acid, 126. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcac catechu, 195, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Ariticial silk, 143. Arum italicum, 106. Assafoetida, 202. Asparagus officinale, 121, 188, 226. Aspartia caid, 315. Asparagus, 295. Asparala odorata, 193. Asphodelus, 115. Asprarguls, 265. Ascarca sativa, 45, 236. Barley, 68, 71, 74, 121, 122, 276, 306, 284 safely grain oil, 20. Bassoric acid, 126. Basevica caid, 126. Basch-nut oil, 20, 32. Beech-nut oil, 20, 32. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beecho-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 279, 347. Beech-nut oil, 20, 32. Beetroot, 68, 74, 128, 252, 27, 279, 28, 281. Beetroot, 68, 74, 128, 252,		
344, 346, 369.		
246, 256. Anthoxanthin, 214, 247. Anthriscus exerfolium, 385. — vulgaris, 288. Anti-enzymes, 358. Anti-enzymes, 366. Aphis chinensis, 212. Apigenin, 246. Apium Petroseliuum, 246. Apoonraphine, 269. Apple, 68, 121, 128. — seed oil, 20. Apposition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabinon-trigalactan-geddic acid, 126. Bean, 132, 371. Beech-nut oil, 20, 32. Beeswax, 43. Beetroot, 68, 74, 128, 255, 289; yield of sugar, 69. Begonia, 305. Beneic acid, 7. Benedict's solution, 85. Benzaldehyde, 181, 182. Benenic acid, 7. Benedict's solution, 85. Benzaldehyde, 181, 182. Beneicatic solution, 85. Berberine, 269. Berberine, 269. Begonia, 305. Bethoric aid, 7. Benedict's solution, 85. Berberine, 269. Berberine, 269. Begonia, 305. Bethoric aid, 7. Benedict's solution, 85. Berberine, 269. Berberine, 269. Begonia, 305. Bethoric aid, 7. Benedict's solution, 85. Berberine, 269. Berberine, 269. Begonia, 305. Bethoric aid, 7. Benedict's solution, 85. Berberine, 269. B		
Anthocyanins and anthocyanthins, 251. Anthorosthin, 244, 247. Anthriscns eerefolium, 385. — vulgaris, 288. Anti-enzymes, 358. Anti-enzymes, 358. Anti-enzymes, 358. Anti-enzymes, 366. Afhis chinensis, 212. Apigenin, 246. Apour Petroselinum, 246. Apocynacæ, 205. Apomorphine, 269. Apposition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, Arabin, 124, 125. Arabin, 124, 125. Arabinon-urgalactan-geddic acid, 126. Arabinsee, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Arachidic acid, 7, 20. Araliacæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcga catechu, 195, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asparagus, 295. Aspardus officinale, 121, 188, 226. Astornal advarata, 193. Asphadelus, 115. Asparagus, 205. Astornal advarata, 193. Asphadelus, 115. Asparagus, 205. Astornal advarata, 193. Asphadelus, 115. Asparagus, 205. Astornal advarata, 193. Asphadelus, 115. Astornal advarata, 193. Asparagus, 205. Astornal advarata, 193. Asphadelus, 115.		324, 346, 369.
Anthoxanthin, 244, 247. Anthrisras exerefolium, 385. — vnlgaris, 288. Anti-enzymes, 356. Antiseptics and enzymes, 366. Aphis chinensis, 212. Apgenin, 246. Apocynacæ, 205. Appomphine, 256. Appomphine, 256. Apposition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabin acid, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Arachis, germination, 41. Arachis oil, 19, 22. Arachis, germination, 41. Arachis oil, 19, 22. Arachidic acid, 7, 20. Aractic plants, 115, 255. Arctic plants, 1		Barley grain oil, 20.
Authriscus eerefolium, 385. — vulgaris, 288. Anti-enzymes, 358. Anti-enzymes, 366. Aphis chieves, 366. Apposition author, 240. Apis prefered of suparticle of s		
- vulgaris, 288. Antiseptics and enzymes, 366. Aphis chinensis, 212. Apigenin, 246. Apocynaceæ, 265. Apomorphine, 269. Appel, 68, 121, 128 seed oil, 20. Apposition theory, 101. Apricot, 70 kernel oil, 19. Aquatic plants, 115, 127, 128, 255, Arabinon-trigalactan-geddic acid, 126. Arabinon-trigalactan-geddic acid, 126. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Aractic plants, 115, 255. Arcca catechu, 195, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoctida, 202. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asporgillus, 73, 215. Astriduum, 194, 236. Astragalus, 126. Atropne, 367. Aucuea sativa, 45, 236. Beestroot, 68, 74, 128, 252, 279, 347. 359; yield of sugar, 69. Begonia, 305. Benezic's solution, 85. Benezideryade, 181, 182. Benezid cxid, 7. Benedict's solution, 85. Berezida, 305. Berezida, 305. Berezida, 305. Berezida, 496. Bethenic acid, 7. Benedict's solution, 85. Beteriot, 68, 74, 128, 252, 279, 347. Beetroot, 68, 74, 128, 252, 279, 347. Beetroot, 68, 74, 128, 252, 279, 347. Beetnot, 68, 74, 128, 255, 359; yield of sugar, 69. Begonia, 305. Berezida, 470. Benedict's solution, 85. Berezida, 470. Benedict's solution, 85. Berezida, 49. Berberic acid, 7. Benedict's solution, 85. Berzolade, 182. Benzulade, 184. Bethenic acid, 7. Benedict's solution, 85. Berezida, 49. Brichletia, 301. - excelsa, 323. Bettoue, 68, 74, 128, 255, Betula acid, 7. Betula alba, 385. Biasting gelatine, 142. Boiled oil, 4. Boragia ve beetroot. Betavulgaris v. beetroot. Betavulgaris v. beetroot. Betavulgaris v. beetroot. Betavulgaris v. beetroo		Dean, 132, 371.
Anti-enzymes, 358. Anti-enzymes, 366. Aphis chinensis, 212. Apigenin, 246. Apour petroselinum, 246. Apocynacæ, 265. Apomorphine, 269. Apple, 68, 121, 128. — seed oil, 20. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 334. Arabic acid, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliacæ, 183. Arbutin, 172, 174, 202. Arctostaphylos Uva-ursi, 205, 246. Areca catechu, 195, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Arolideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Aspergillus, 73, 215. — niger, 354, 382. Asphodalus, 115. Astridum, 194, 236. Astragalus, 126. Atropne, 367. Aucota paponica, 199. Autotermentation, 376. Avena sativa, 45, 236. Bectroot, 68, 74, 128, 252, 279, 347. 359; yield of sugar, 69. Bemclict's solution, 85. Benealct's solution, 85. Beneacic's solution, 85. Beneacic's solution, 85. Benedict's solution, 85. Benealct's solution, 85. Benealct's solution, 85. Bereiot's Solution, 95. Benedict's solution, 95. Benedict's solution, 95. Benedic's solution, 95. Benedic's solution, 95. Benedic's solution, 95. Berediot's Solution, 95. Bereiot's Solution, 95. Benedic's Solution, 95. Benedic's Solution, 95. Benedic's Solution, 95. Benedic's solution, 95. Bereiot's Solution, 95. Beneziclehyde, 181, 182. Benzyl alcohol, 386. Berbroot, 96. Bertroot, 76. Bettavulgaris v. beetroot. Betaine, 47, 275. Betta vulgaris v. beatine, 147. Betaine, 147, 275. Betta vulgaris v. beatine, 147. Betaine, 147, 275. Betta vulgaris v. beatine, 147. Betaine, 147,		
Antiseptics and enzymes, 366. Aphise hinensis, 212. Apigenin, 246. Apium Petroselinum, 246. Apocymaceæ, 205. Apomorphine, 269. Apple, 68, 121, 128. — seed oil, 20. Apposition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinon-trigalactan-geddic acid, 126. Arachidic acid, 7, 20. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcea catechu, 195, 266, 275. Arceoline, 266. Arcea catechu, 195, 266, 275. Arceoline, 266. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asphorellhus, 73, 215. Asphorellus, 73, 215. Asphorelus, 74, 202. Area oil oil, 4, 203. Benedict's solution, 85. Benenic acid, 7, 2 Benedict's solution, 85. Benenic acid, 7, 2 Benezide, 74, 275. Benedict's solution, 85. Benezide, 74, 275. Behenic acid, 7, 2 Bertholletia, 301. — excelsa, 323. Betula alba, 385. Bilberry, 249. Birch, 68. Birute reaction, 396. Boirtus, 390. Boletus adulas, 132. Blasting gelatine, 142. Boiled oil, 4. Bo		
Aphis chinensis, 212. Apigenin, 246. Apium Petroselinum, 246. Apomorphine, 269. Apple, 68, 121, 128. — seed oil, 20. Apposition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachisic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Arachis, germination, 41. Boiled oil, 4. Boiled oil, 4. Boiled oil, 4. Boraginaceæ, 195, 222, 396. Borneol, 386. Brossica Napus, 2, 4, 340. — Rapa, 4. Brassidic acid, 7, Brazil nut, amount of fat, 2. Bromelin, 348, 353. Brownel, 386. Bryonol, 386. Brozilachyde, 181, 182. Benezic acid, 7. Bethollctia, 301. — excelsa, 323. Beta vulgaris v. beetroot. Betaine, 47, 275. Betula alba, 385. Bilberry, 249. Birch, 68. Biuret: caction, 30. Bettworling arbic vice derivatives of proteins, 313. Blasting gelatine, 142. Boiled oil, 4. Boiled oil, 4. Boraginaceæ, 195, 222, 396. Borreol, 386. Bryonol, 386. Bryonol, 386. Bryonol, 386. Bryonol, 386. Bryonol, 386. Brozil albey, 249. Brich, 68. Biuret caction, 30. Betta vulgaris v. beetroot. Betaine, 47, 275. Betula alba, 385. Biuretic derivatives of proteins, 313. Blasting gelatine, 142. Boiled oil, 4. Boiled oil, 4. Boraginaceæ, 195, 222, 396. Broreol, 386. Bryonol, 386. Broreira addicacin, 102. Cacho butter, 2, 3, 32. Cadaverine, 274. Cacho butter, 2, 3, 32. Cadaverine, 274. Cacho butter, 2, 3, 32. Cadaverine, 277. Cadreine,		252, 279, 347,
Apigenin, 246. Apocynaceæ, 265. Apomorphine, 260. Apple, 68, 121, 128. — seed oil, 20. Appicot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabinon-trigalactan-geddic acid, 126. Arabinon-trigalactan-geddic acid, 126. Arabinos, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asparagus, 295. Asphodellus, 115. Astidium, 194, 236. Astropala, 360. Benealcit's solution, 85. Benealcit's solution, 386. Berberine, 269. Bertholdetia, 301. —excelsa, 323. Beta vulgaris v. beetroot. Betaine, 47, 275. Beta vulgaris v. beetroot. Betaine, 47, 275. Beta vulgaris v. beatroot. Betaine alac, 47, 275. Beta vulgaris v. beatroot. Betaine, 47, 275. Beta vulgaris v. beatroot. Betaine, 47, 275. Bitertyne, 27, 278. Biberry, 249. Birch, 68. Birch, 68. Birch of setula alac, 172. Birch, 68. Birch of setula alac, 172. Boracion, 370. Bircholetia, 300. Boletis siquita		Begavia 205
Apium Petrosclinum, 246. Apocynaceæ, 265. Apomorphine, 269. Apple, 68, 121, 128. — seed oil, 20. Approsition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, Arabin 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Aratic plants, 115, 255. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Arioticke, 114. Artificial silk, 143. Arum italicum, 106. Asafoctida, 202. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asporgillus, 73, 215. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparadus, 126. Astrogalus, 126. Astrogalus, 126. Atropine, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Benezdict's solution, 85. Benezaldeyde, 181, 182. Benzaldeyde, 181, 182. Benzyl alcohol, 386. Berberine, 269. Bethoiletia, 301. — excelsa, 323. Betula alba, 385. Bilberry, 249. Birch, 68. Biuret reaction, 307. Biuretic derivatives of proteins, 313. Blasting gelatine, 142. Blown oil, 4. Boiled oil, 4. Boiled oil, 4. Boiled oil, 4. Boiled oil, 4. Boragineaeæ, 195, 222, 396. Borneol, 386. Bryunoil, 14. Boragiaria, 194, 25. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Brustal'is experiment, 14. Cacab butter, 2, 3, 32. Cadaverine, 274. Cactus, 126. Cadaverine, 274. Cardeine, 277, 278, 281. Caffeine, 277, 278, 281. Caffeine, 277, 278, 281. Calanthe, 190. Cardenia dava 49. Birch, 68. Bretdil alba, 385. Biasting gelatine, 142. Boiled oil, 4. Boiled oil, 4. Boragiariaeæ, 195, 222, 396. Borneol, 386. Bryunol, 184, 266. Bryunol, 184, 266. Bryunol, 184, 266. Bryund 1		Behenic acid 7
Apocynacæ, 265, Apomorphine, 269, Apple, 68, 121, 128, — seed oil, 20. Apposition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, Arabin, 124, 125. Arabin, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183, Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Ura-ursi, 205, 246. Arcea catechu, 195, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Arini talicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Asparagine, 315. Aspergillus, 73, 215. — niger, 354, 382. Asperula odorata, 193. Asphadelus, 115. Astridum, 194, 236. Astragalus, 126. Atropine, 367. Aucota paponica, 199. Aucotermentation, 376. Avena sativa, 45, 236. Benzaldehyde, 181, 182. Benzyl alcohol, 386. Berberine, 269. Bertholletia, 301. —excelsa, 323. Beta vulgaris v. beetroot. Betaine, 47, 275. Betula alba, 385. Bilberry, 249. Bilberry, 249. Bilberty, 249. Bilberty, 249. Bilberth, 261. Betavulgaris v. beetroot. Betaine, 47, 275. Betula alba, 385. Bilbery, 249. Bilberth, 269. Bettholletia, 301. —exclsa, 323. Beta vulgaris v. beetroot. Betaine, 47, 275. Betula alba, 385. Bilbery, 249. Biloretic derivatives of proteins, 313. Blasting gelatine, 142. Boletus, 390. Boletus sdulis, 73, 114. Boletus, 390. Boletus derivatives of proteins, 313. Blasting gelatine, 142. Blown oil, 4. Boletus, 390. Boletus derivatives of proteins, 313. Blasting gelatine, 142. Blown oil, 4. Boletus, 390. Boletus, 390. Brassica Carlothus, 73, 114. Boletus, 390. Boletus derivatives of proteins, 313. Blasting gelatine, 142. Blown oil, 4. Boletus, 390. Brassica Carlothus, 73, 114. Boletus, 390. Brassica Carlothus, 72, 4, 340. — clreacea, 340. — Rapa,		
Appmorphine, 269. Apple, 68, 121, 128. — seed oil, 20. Apposition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcea catechin, 195, 266, 275. Arecoline, 266. Arcea catechin, 195, 266, 275. Arecoline, 266. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asphrogallus, 73, 215. Casparagine, 315. Asphrogallus, 73, 215. Asphrogallus, 73, 215. Asphrogans officinale, 121, 188, 226. Aspartic acid, 315. Asphrogallus, 73, 215. Cactus, 126. Cadaverine, 274. Cacavalpinia, 194. Cacavalpini		
Apple, 68, 121, 128. — seed oil, 20. Appricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachisic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Arachis, germination, 41. Arachis oil, 19, 22. Arachisin, 172, 174, 202. Articolants, 115, 255. Arctostaphylos Uva-uvsi, 205, 246. Areca catechu, 195, 266, 275. Arecoline, 266. Argnine, 316; occurrence of, in plant, 321, 328. Aroideze, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asphodelus, 115. Asphodelus, 115. Asphodelus, 115. Asphodelus, 115. Asphodelus, 115. Astidium, 194, 236. Astrogalus, 126. Autopene, 367. Autocum, 367. Au		
seed oil, 20. Apposition theory, 101. Apricot, 70 kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinon-trigalactan-geddic acid, 126. Arabinos, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Argenine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315		
Approsition theory, 101. Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachis, germination, 41. Arachis oil, 19, 22. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Aractic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Argenine, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoctida, 202. Asparagus, 205. Asparatic acid, 315. Asparagus, 205. Asparatic acid, 315. Asperillus, 73, 215. Asperillus, 73, 215. Asperillus, 73, 215. Asphodelus, 115. Astidium, 194, 236. Astrogalus, 126. Atropne, 367. Aucota planta, 193. Asphodelus, 115. Astrogalus, 126. Atropne, 367. Aucota planta, 193. Asphodelus, 115. Astorium, 194, 236. Astrogalus, 126. Atropne, 367. Aucota planta, 193. Asphodelus, 115. Cacatpinica, 194. Cacatpinica, 195. Calabar bean, 385. Betal vulgaris v. beetroot. Betaine, 47, 275. Betula alba, 385. Bisting 7249. Birch, 68. Bisuret reaction, 307. Biuretic derivatives of proteins, 313. Blasting gelatine, 142. Boiled oil, 4. Boiled oil, 4. Borellus, 73, 114. Boraginacee, 195, 222, 396. Brassidic acid, 7, 30. Brassidic acid, 7, 36. Brassidic acid, 7, 36. Bryonol, 386. Cadaverine, 27, 278, 281. Cadiene, 272, 278, 281. Caffeine, 272, 278, 281. Cacatpinica, 142. Boiled oil, 4. Boiled oil, 4. Borelius, 390. Boletus, 390. Boletus advais, 112. Calcan acid, 134. Borelius, 182. Borelius, 183. Blasting 249. Birch, 68. Brotal alba, 185. Basting 249. Birch, 68. Brotal alba, 185. Calcatus, 193. Calcatus, 193. Cacato butter, 2, 3, 32. Cadaverine, 274. Cacato butter, 2, 3, 32. Cadaverine, 274. Cacato but		
Apricot, 70. — kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabin, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabiniose, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Argeinie, 316; occurrence of, in plant, 321, 328. Aroiteæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Asparerillus, 73, 215. — niger, 354, 382. Asperula odorata, 193. Asphadelus, 115. Astridum, 194, 236. Astragalus, 126. Atropne, 367. Aucote acterin, 199. Aucotermentation, 376. Avena sativa, 45, 236. Betal vulgaris v. beetroot. Betaine, 47, 275. Betula alba, 385. Bilberry, 249. Bilberry, 249. Bilberty, 249. Bilberty, 249. Bilberty, 249. Bilbert, 80. Biuretic derivatives of proteins, 313. Blasting gelatine, 142. Booled oil, 4. Boietus, 390. Boletous derivatives, 390. Boletous derivatives of proteins, 313. Blasting gelatine, 142. Blown oil, 4. Boietus, 390. Boletous, 390. Boletous, 390. Boletous, 47. Boraginaceæ, 195, 222, 396. Borneol, 386. Brassicia caid, 7. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Brucine, 26. Boletous, 390. Boletous, 300. Brassica caid, 7. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Brucine, 26. Caracobiute, 121. Boraginaceæ, 195, 222, 396. Borneol, 386. Brassica caid, 7. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Brucine, 26. Caracobiute, 122. Blown oil, 4. Boletous derivitives of proteins, 313. Blasting gelatine, 142. Blown oil, 4. Boletous derivitives of proteins, 313. Blasting gelatine, 142. Boletous derivitée derivatives of proteins, 313.		
— kernel oil, 19. Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabin, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcea catechu, 195, 266, 275. Arecoline, 266. Areginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Aruni italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asprorgillus, 73, 215. Cacao butter, 2, 3, 32. Cactus, 126. Cadaverine, 274. Cacaslpinia, 194. Cacaslpinia, 194. Cacaslpinia, 194. Cacaslpinia, 194. Cacaslpinia, 194. Cacalanthe, 195. Calabar bean, 386. Bilberry, 249. Birch, 68. Bivert eaction, 307. Biasting gelatine, 142. Boiled oil, 4. Boile		
Aquatic plants, 115, 127, 128, 255, 334. Arabic acid, 124, 125. Arabin, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabiniose, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Arctic plants, 115, 255. Arteotaphylos Uva-ursi, 205, 246. Arcea catechu, 195, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagus, 295. Asparagus officinale, 121, 188, 226. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus officinale, 121, 188, 226. Aspareillus, 73, 215. — niger, 354, 382. Asphodelus, 115. Astidium, 194, 236. Astragalus, 126. Atropine, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Bilberry, 249. Bitch, 68. Biuret caction, 307. Biureti derivatives of proteins, 313. Blasting gelatine, 142. Blown oil, 4. Boletus, 390. Boletus, 300. Bortel, 386. Borneol, 386. Borneol, 386. Bryonol, 386. Cactus, 126. Cactus, 126. Cadavetine, 274. Caralpinia, 194. — coriaria, 194, 205, 207. Callanthe, 191. Campanula ceae, 115. Campanula trachclium, 301. Canalgre, 195, 210, 211. Cananga odorata, 385.		
334. Arabin (124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Argenine, 316; occurrence of, in plant, 321, 328. Arolideæ, 176. Artichoke, 114. Artificial silk, 143. Artuficial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asprogallus, 73, 215. — niger, 354, 382. Asphordelus, 115. Astidium, 194, 236. Astrogalus, 126. Attopine, 367. Autotermentation, 376. Avena sativa, 45, 236. Backleus macerans, 103. Bilberry, 249. Bilberty, 249. Biuret reaction, 307. Biusetic derivatives of proteins, 313. Biusting gelatine, 142. Boiled oil, 4. Boiled oi		
Arabic acid, 124, 125. Arabin, 124, 125. Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachis germination, 41. Arachis oil, 19, 22. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Argeinine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoctida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 205. Asparatic acid, 315. Asparatic acid, 315. Asperillus, 73, 215. Asperillus, 73, 215. Asperillus, 73, 215. Asphodelus, 115. Asphodelus, 115. Astidum, 194, 236. Astrogalus, 126. Attopine, 367. Aucuta japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Birch, 88. Biuret reaction, 307. Biuretic derivatives of proteins, 313. Blasting gelatine, 142. Boiled oil, 4. Boiled oil, 4. Boiled oil, 4. Boraginaceæ, 195, 222, 396. Brassica Napus, 24, 340. — eliveacea, 340. — Rapa, 4. Brassilic acid, 7. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Bryonol, 386. Cackao butter, 2, 3, 32. Cactus, 126. Cadaverine, 274. Cactus, 126. Cadaverine, 274. Cactus, 126. Cadaverine, 274. Cactaslpinia, 194. Carcialine, 142. Boiled oil, 4. Boiled oil, 4. Bordetus, 390. Bortins derivatives of proteins, 313. Blasting gelatine, 142. Boiled oil, 4. Bordetus, 390. Boretus advis, 5, 306. Bryonol, 386. Bryonol, 386. Bryonol, 386. Bryonol, 386. Bryonol, 386. Cacaco butter, 2, 3, 32. Cacaths, 126. Cadaverine, 274. Cacaslpinia, 194. — coriaria, 194, 205, 207. Caffeine, 277, 278, 281. Campanula trachclium, 301. Canapanula trachclium, 301. Canaligre, 195, 210, 211. Cannagna odorata, 385. Candolleaceæ, 115.		
Arabin, 124, 125, Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129, Arachidic acid, 7, 20. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Arachis, germination, 41. Arachis oil, 19, 22. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcea catechu, 195, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Arbitichoke, 114. Artificial silk, 143. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus officinale, 121, 188, 226. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus officinale, 121, 188, 226. Asparenta acid, 315. Aspervillant, 73, 215. — niger, 354, 382. Asphodelus, 115, Astidium, 194, 236. Astragalus, 126. Atropne, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Biureti cedrivatives of proteins, 313. Blasting gelatine, 142. Blown oil, 4. Boietus, 390. Boletus, 49, 242, 396. Borneol, 386. Borneol, 386. Borneol, 386. Brassical value, -195, 222, 396. Borneol, 386. Brassical caid, 7. Brassidic acid, 7. Brazil nut, amount of fat, 2. Broatine, 172. Blown oil, 4. Boletus, 390. Boletus ednlis, 73, 114. Boraginacee, 195, 222, 396. Borneol, 386. Brassica caid, 7. Brazil nut, amount of fat, 2. Cacho butter, 2, 3, 32. Cacho butter, 2,		
Arabinon-trigalactan-geddic acid, 126. Arabinose, 53, 59, 126, 129. Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 205. Asparagine, 205. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagos officinale, 121, 188, 226. Aspartic acid, 315. Asphorgillus, 73, 215. — niger, 354, 382. Asphodelus, 115. Astidium, 194, 236. Astropala, 195. Astropane, 367. Autotermentation, 376. Avena sativa, 45, 236. Buretic derivatives of proteins, 313. Blasting gelatine, 142. Boiled oil, 4. Boile oil, 4. Boiled oil, 4. Boiled oil, 4. Boiled oil, 4. Bola biel o		
Arachidic acid, 7, 20. Arachis, germination, 41. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcacine, 266. Arginine, 316; occurrence of, in plant, 321, 328. Artichoke, 114. Artificial silk, 143. Artificial silk, 143. Artinicial silk, 143. Brucine, 268. Bryonol, 386. Bryonol, 386. Bryonol, 386. Bryonol, 386. Bryony, 386. Cactus, 126. Cactus, 126. Cadaverine, 274. Casalpinia, 194. Cardiene, 277, 278, 281. Caffeine, 277, 278, 281. Caffeine, 277, 278, 281. Caffeine, 277, 278, 281. Campanula ceae, 115. Campanula trachclium, 301. Canagire, 195, 210, 211. Canagire, 195, 210, 211. Canagire, 195, 210, 211.	Arabinon-trigalactan-geddic acid, 126.	Biuretic derivatives of proteins, 313.
Arachis, germination, 41. Arachis oil, 19, 22. Arachis oil, 19, 22. Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arganine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Artim italicum, 106. Asafoctida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 205. Asparatic acid, 315. Asparatic acid, 315. Asperillus, 73, 215. Asperillus, 73, 215. Asperillus, 73, 215. Asphodelus, 115. Asphodelus, 115. Astidium, 194, 236. Astrogalus, 126. Attopine, 367. Autotermentation, 376. Avena sativa, 45, 236. Boiletus, 390. Botchis caplis, 52, 222, 396. Brassica Napus, 2, 4, 340. —— colcreacea, 340. —— Rapa, 4. Brassilic acid, 7. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Bryonol, 386. Cactos butter, 2, 3, 32. Cactus, 126. Cadaverine, 274. Cactus, 126. Cadaverine, 274. Cactus, 126. Cadeverine, 274. Cactus, 126. Cadeverine, 274. Cactus, 126. Cadaverine, 274. Cactus, 126. Cadaverine, 274. Cactus, 126. Cadaverine, 274. Cactus, 126. Cadaverine, 277. Caffeine, 277, 278, 281. Caffetamic acid, 193. Calabar bean, 386; fat of, 17. Campanula trachclium, 301. Canagine, 195, 202, 207. Caffeine, 290.	Arabinose, 53, 59, 126, 129.	Blasting gelatine, 142.
Arachis oil, 19, 22. Arachis oil, 19, 21, 25. Arachis oil, 19, 22. Arachis oil, 19, 22. Boletus edulis, 73, 114. Boraginacee, 195, 222, 396. Borneol, 386. Borneol, 386. Brassiale acid, 7. Brasil out, anount of fat, 2. Brazil nut, amount of fat, 2. Brazil nut, am	Arachidic acid, 7, 20.	Blown oil, 4.
Araliaceæ, 183. Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcca catechu, 195, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Aruni italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asprorgillus, 73, 215. — niger, 354, 382. Asphordalodus, 115. Astridium, 194, 236. Astropalus, 126. Attopine, 367. Aucuta japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Borneol, 386. Brassida Napus, 24, 340. — clreacea, 340. — Rapa, 4. Brassidic acid, 7. Brazil nut, amount of fat, 2. Bromellin, 348, 353. Bryonol, 386. Carcal initial, 202. Cactus, 117. Cacatenin, 194. Cacatenin, 194. Cacatenin, 194. Carcal initial, 194. — coriaria, 194, 205, 207. Callanthe, 191. Campanula trachelium, 301. Canaigre, 195, 210, 211. Cananga odorata, 385.	Arachis, germination, 41.	Boiled oil, 4.
Arbutin, 172, 174, 202. Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcea catechu, 195, 266, 275. Aregoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artichoke, 114. Artichoke, 114. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus officinale, 121, 188, 226. Asperula odorata, 193. Asphodelius, 115. Astidium, 194, 236. Astragalus, 126. Atropine, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Boraginaceæ, 195, 222, 396. Brassica Napus, 2, 4, 340. — akapa, 4. Brassidic acid, 7. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Brucine, 268. Bryonol, 386. Bryonol, 386. Bryony, 386. Bryony, 386. Bryony, 386. Bryony, 386. Cacha butter, 2, 3, 32. Cacha butter, 2, 3, 32. Cachaverine, 274. Cesalpinia, 194. — coriaria, 194, 205, 207. Callabar bean, 386; fat of, 17. Calampanula trachclium, 301. Canaigre, 195, 210, 211. Cananga odorata, 385. Calabal bean, 386; fat of, 17. Campanula trachclium, 301. Canaigre, 195, 210, 211. Cananga odorata, 385.		Boletus, 390.
Arctic plants, 115, 255. Arctostaphylos Uva-ursi, 205, 246. Arcea catechu, 195, 266, 275. Aregoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Artinicial silk, 143. Artinicial silk, 143. Artinicial silk, 143. Artinicial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Aspergillus, 73, 215. — niger, 354, 382. Asphodelus, 115. Astidium, 194, 236. Astragalus, 126. Attropne, 367. Aucotermentation, 376. Avena sativa, 45, 236. Borneol, 386. Brassica Napus, 2, 4, 340. — cleracea, 340. — Rapa, 4. Brassidic acid, 7. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Brucine, 268. Bryonol, 386. Bryonol, 386. Bryonol, 386. Bryonol, 386. Carcum, 117. Butea, 201. Bütschli's experiment, 14. Cacadaptine, 274. Cacadaverine, 274. Cacadaverine, 274. Casadpinia, 194. — coriaria, 194, 205, 207. Callanthe, 191. Campanula creachium, 301. Canaigre, 195, 210, 211. Cananga odorata, 385. Candolleaceæ, 115.		Boletus edulis, 73, 114.
Arctos Infly los Uva-ursi, 205, 246. Arcea catechu, 195, 266, 275. Arceoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Aspartic acid, 315. Cactos butter, 2, 3, 32. Cactus, 126. Cadaverine, 274. Cactus, 126. Cadverine, 274. Cactos juiia, 194. — coriaria, 194, 205, 207. Caffeine, 277, 278, 281. Caffetamic acid, 193. Calabar bean, 386; fat of, 17. Campanula trachclium, 301. Campanula trachclium, 301. Canaigre, 195, 210, 211. Cananga odorata, 385. Candolleaceæ, 115.		Boraginaceæ, 195, 222, 396.
Areca catechu, 195, 266, 275. Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artichoke, 114. Artincial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus officinale, 121, 188, 226. Asparagins officinale, 121, 188, 226. Asparegillus, 73, 215. — niger, 354, 382. Asphoellus, 115. Astidium, 194, 236. Astragalus, 126. Atropine, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Bacillus macerans, 103. — olcraeca, 340. — Rapa, 4. Brassidic acid, 7. Brazil nut, amount of fat, 2. Brazil nut, amount of autoric, particular, particul		Borneol, 386.
Arecoline, 266. Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Aruni italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asporgillus, 73, 215. — niger, 354, 382. Asphordellus, 115. Asphordellus, 115. Astidium, 194, 236. Astrogalus, 126. Attopine, 367. Autotermentation, 376. Avena sativa, 45, 236. Bacillus macerans, 103. Are Rafa, 4. Brassidic acid, 7. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Bryunn, 17. Butea, 201. Bryum, 17. Butea, 201. Caca, 201. Caca, 201. Caca, 201. Caca, 201. Caca, 201. Cacatus, 126. Cacatus, 126. Cadaverine, 274. Cacastlinia, 194. — coriaria, 194, 205, 207. Callanthe, 191. Campanula trachelium, 361. Canagine, 195, 210, 211. Canagine, 195, 210, 211. Canagine, 275.		Brassica Napus, 2, 4, 340.
Arginine, 316; occurrence of, in plant, 321, 328. Aroideæ, 176. Artichoke, 114. Artichoke, 114. Artichoke, 114. Asparagine, 315. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asperula odorata, 193. Asphodelius, 115. Astridium, 194, 236. Astragalus, 126. Atropine, 367. Aucueba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Backleus macerans, 103. Brassidic acid, 7. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Brucine, 268. Bryonot, 386. Bryony, 386. Cackao butter, 2, 3, 32. Cactus, 126. Cactus, 126. Cackao butter, 2, 3, 32. Cactus, 126. Cactus, 126. Cackao butter, 2, 3, 32. Cactus, 126. Cactus (14, 2. Bromelin, 348, 353. Brucine, 268. Bryonot, 386. Bryony, 386. Bryond, 386. Bryond, 386. Bryony, 386. Broon, 268. Cactus (148. Cactus (148. Cactus (148. Cactus (148. Cactus (148. Cactus (148	Arcea catechu, 195, 266, 275.	
321, 328. Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Artinicial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagos officinale, 121, 188, 226. Aspartic acid, 315. Aspergillus, 73, 215. — niger, 354, 382. Asphrola odorata, 193. Asphodelus, 115. Astidium, 194, 236. Astrogalus, 126. Atropne, 367. Aucotementation, 376. Avena sativa, 45, 236. Brazil nut, amount of fat, 2. Bromelin, 348, 353. Brucine, 268. Bryono, 386. Bryono, 386. Bryono, 186. Asynm, 117. Butea, 201. Butechli's experiment, 14. Cacton butter, 2, 3, 32. Cactus, 126. Cadaverine, 274. Casalpinia, 194. — coriaria, 194, 205, 207. Caffeine, 277, 278, 281. Caffetannic acid, 193. Calabar bean, 386; fat of, 17. Calanthe, 191. Campanula trachelium, 301. Canagre, 195, 210, 211. Canagra odorata, 385. BACILLUS macerans, 103.		
Aroideæ, 176. Artichoke, 114. Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Aspartic acid, 315. Aspartic acid, 315. Aspartic acid, 315. Asperillus, 73, 215. — niger, 354, 382. Aspfordla odurata, 193. Asphodelus, 115, Astidium, 194, 236. Astrogalus, 126. Atropine, 367. Autotementation, 376. Avena sativa, 45, 236. Bromelin, 348, 353. Bruone, 268. Bryono, 386. Cactan it, 20. Cactan it, 20. Cadaverine, 27, 21, 23. Cadaverine, 274. Cardsplinia, 194. — coriaria, 194, 205, 207. Caffeine, 277, 278, 281. Caffetamic acid, 193. Calabar bean, 386; fat of, 17. Campanula trachelium, 301. Canagie, 195, 210, 211. Cananga odorata, 385. Candolleaceæ, 115.		
Artichoke, 114. Bryonol, 386. Bryony, 386. Bryond, 286. Bryony, 386. Bryond, 286. Bryond, 286. Bryond, 286. Bryond, 286. Bryony, 386. Bryond, 286. Bryond, 296. Asparable, 21. Cachus,		Brazil nut, amount of fat, 2.
Artificial silk, 143. Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Aspartic acid, 315. Asperillus, 73, 215. — niger, 354, 382. Asphordlolus, 115. Astidium, 194, 236. Astragalus, 126. Astrogne, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Backleus Bryonol, 386. Bryandle star, 36. Cachavella, 194. Cachabella, 115. Cachabella, 115. Cachabel		Bromelin, 348, 353.
Arum italicum, 106. Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 205. Asparagns officinale, 121, 188, 226. Asparegus, 205. Asparegulus, 205. Aspergillus, 73, 215. — niger, 354, 382. Asperula odorata, 193. Asphodelus, 115. Astidium, 194, 236. Astropine, 367. Aucota japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Bryum, 117. Butea, 201. Caccao butter, 2, 3, 32. Cactus, 126. Cacaba tive, 124. Casalpinia, 194. — coriaria, 194, 205, 207. Caffetannic acid, 193. Calbara bean, 386 ; fat of, 17. Campanula trachelium, 301. Canaigre, 195, 210, 211. Canang odorata, 385. Candolleaceæ, 115.		
Asafoetida, 202. Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Aspergillus, 73, 215. — niger, 354, 382. Asphodolus, 115. Astidium, 194, 236. Astragalus, 126. Astropne, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Backleus Market and Stryum, 117. Butta, 201. Butta, 201		Bryonoi, 386.
Asparagine, 315. — synthesis of, in plant, 335, 336, 337. Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Aspergillus, 73, 215. — niger, 354, 382. Asphodelus, 115. Astridium, 194, 236. Astropine, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Bitta, 207. Bittea, 207. Cactus, 126. Casalpinia, 194.		
— synthesis of, in plant, 335, 336, 337. Asparagus, 295. Aspartic acid, 315. Aspergillus, 73, 215. — niger, 354, 382. Asphordle odorata, 193. Asphodelus, 115. Astidium, 194, 236. Astragalus, 126. Atropine, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. Backleus macerans, 103. Backleus macerans, 103. Backleus macerans, 103. Bütschli's experiment, 14. Acaco butter, 2, 3, 32. Cachas, 126. Cachas, 126. Cachas, 126. Cachas lean, 194, 205, 207. Caffeine, 277, 278, 281. Caffetannic acid, 193. Calabar bean, 386; fat of, 17. Campanula trachclium, 301. Canaigre, 195, 210, 211. Canaigre, 195, 210, 211. Canadolleaceæ, 115.		
Asparagus, 295. Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Asprogillus, 73, 215. Cactus, 126. Cadverine, 274. Cactus, 126. Cactus, 126. Cactus, 126. Cactus, 126. Cactus, 126. Cactus, 126. Cadverine, 274. Cactus, 126. Ca	- cynthesis of in plant 225 226 227	
Asparagus officinale, 121, 188, 226. Aspartic acid, 315. Aspergillus, 73, 215. — niger, 354, 382. Asphodelus, 115. Astridium, 194, 236. Astropine, 367. Autopine, 367. Automentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Cacatos putter, 2, 3, 32. Cactus, 126. Cadaverine, 274. Casalpinia, 194. Casalpinia, 194. — coriaria, 194, 205, 207. Caffetannic acid, 193. Caffetannic acid, 193. Callabar bean, 386 if at of, 17. Callabar bean, 386 if at of, 17. Campanulacea, 115. Campanula trachelium, 301. Canaigre, 195, 210, 211. Cananga odorata, 385. Candolleaceæ, 115.		Butsein s experiment, 14.
Aspartic acid, 315. Aspergillus, 73, 215. — niger, 354, 382. Asphoellus, 115. Astridium, 194, 236. Astragalus, 126. Atropne, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Cadaverine, 274. Casvalpinia, 194, 205, 207. Castleine, 277, 278, 281. Caffetannic acid, 193. Callabar bean, 386; fat of, 17. Campanula creae, 115. Canagre, 195, 210, 211. Canagra odorata, 385. Candolleaceæ, 115.		Cacao butter 2 2 22
Aspergillus, 73, 215. — niger, 354, 382. — sperula odorata, 193. Asphodelus, 115. Astidium, 194, 236. Astropine, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Cadaverine, 274. Carsalpinia, 194, 205, 207. Caffetine, 277, 278, 281. Caffetannic acid, 193. Callabar bean, 386; fat of, 17. Calanthe, 191. Campanulacea, 115. Canaigre, 195, 210, 211. Canaigre, 195, 210, 211. Canaigre, 195, 210, 211. Canaigre odorata, 385. Candolleaceæ, 115.		
niger, 354, 382. Asproula odorata, 193. Asphodelus, 115. Astidium, 194, 236. Astropine, 367. Autopine, 367. Autobarimentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Casalpinia, 194, 205, 207. Caffetannic acid, 193. Caffetannic acid, 193. Callabar bean, 386; fat of, 17. Callabar bean, 386; fat of, 17. Campanulaceæ, 115. Campanula trachclium, 301. Canaigre, 195, 210, 211. Canagn odorata, 385. Candolleaceæ, 115.		
Aspérula odurala, 193. Asphodelus, 115. Astidium, 194, 236. Astragalus, 126. Astropen, 367. Autopen, 367. Autopenentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. — coriaria, 194, 205, 207. Caffeine, 277, 278, 281. Caffeine, 277, 278, 281. Callabar bean, 386; fat of, 17. Callanthe, 191. Campanulaceæ, 115. Campanula trachclium, 301. Canaigre, 195, 210, 211. Cananga odurala, 385. Candolleaceæ, 115.		
Asphodelus, 115. Astridium, 194, 236. Astropine, 367. Aucuba japonica, 199. Autolermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Caffetannic acid, 193. Callabar bean, 386; fat of, 17. Callanthe, 191. Campanulacea, 115. Campanula (rachelium, 301. Canaigre, 195, 210, 211. Canaigre odorata, 385. Candolleaceæ, 115.		
Astridium, 194, 236. Astragalis, 126. Atropine, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Caffetannic acid, 193. Calabar bean, 386 ; fat of, 17. Campanulaceæ, 115. Campanula trachclium, 301. Canaigre, 195, 210, 211. Canago adorata, 385. Candolleaceæ, 115.		Caffeine, 277, 278, 281.
Astragalus, 126. Atropine, 367. Anacuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Calabar bean, 386; fat of, 17. Calanthe, 191. Campanulaceæ, 115. Campanula trachelium, 301. Canaigre, 195, 210, 211. Canaigre odorata, 385. Candolleaceæ, 115.		Caffetannic acid, 103.
Atropine, 367. Aucuba japonica, 199. Autotermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Calanthe, 191. Campanulacea, 115. Canaigre, 195, 210, 211. Canaigre odorata, 385. Candolleaceæ, 115.		
Aucuba japonica, 199. Autofermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Campanulaceæ, 115. Campanula (rachelium, 301. Canaigre, 195, 210, 211. Cananga odorata, 385. Candolleaceæ, 115.		
Autotermentation, 376. Avena sativa, 45, 236. BACILLUS macerans, 103. Canaga odorata, 385. Candolleaceæ, 115.		
Avena sativa, 45, 236. BACILLUS macerans, 103. Canaigre, 195, 210, 211. Cananga odorata, 385. Candolleaceæ, 115.		
BACILLUS macerans, 103. Cananga odorata, 385. Candolleaceæ, 115.		Canaigre, 195, 210, 211.
BACILLUS macerans, 103. Candolleaceæ, 115.		Cananga odorata, 385.
	BACILLUS macerans, 103.	
		Cane sugar, 68; estimation of, 81;

INDEX

inversion of, 70; manufacture of, 68;	Chicory, 114, 121; see also Cichorium.
properties and reactions of, 70.	Chinese galls, 212.
Canna, 301.	— insect wax, 43.
— indica, 98.	Chlorine and tannin formation, 196.
Cannabis indica, 275.	Chlorophyll, 225, 272; colorometric
— sativa, 275, 323, 371, 372; germina-	estimation of, 239; constitution, 231;
tion, 40.	crystalline and amorphous, 233; ex-
Caprifoliaceæ, 183.	traction, 238; formation, 227; phy-
Caragheen mucilage, 127.	sical properties, 234.
Carbohydrates, classification of, 53.	Chlorophyll and hæmoglobin, 236.
Carica papaya, 348, 370, 372.	Chlorophyllase, 234, 235. Chlorophyllide, 235.
Carnauba wax, 43.	Chlorophyllin, 232, 233.
Carnaubic acid, 43. Carotin, 229, 241, 242, 262.	Chlorophyllogen, 227.
Carotinoids, 229, 241.	Chlorophytum, 225.
Carrot, 68, 128, 242, 273.	Chloroplasts, 2, 24, 35, 191, 225, 241,
Carubin, 121.	301; distribution, 226; of parasites,
Carum carvi, 385.	227; origin, 225; structure, 225.
Caryophyllaceæ, 66, 396.	Cholesterol, 16, 43, 185, 186.
Cascarilla hexandra, 60.	Choline, 47, 274, 275.
Cassia, 211.	Chondriosomes, 226.
- obovata, 128.	Chromogens, 253, 255, 261.
Castanca, 44, 194, 195.	Chromoprotein, 325, 327.
Castor oil, 2, 4, 19, 32, 300, 302,	Chrysin, 245.
367; fat splitting enzyme in, 4, 366,	Chrysophyll, 243.
367.	Cichorium endiva, 340.
Catalase, 161, 349.	Cichorium Intibus, 114.
Calalysis, 345.	Cinchona, 266, 268, 273, 281.
Catechin, 203, 204, 217; tannins, 210.	Cinchonine, 268. Cinnamic alcohol, 386.
Catechol = pyrocatechol, 190, 201. Catecholase, 190.	Citronellol, 386.
Catechu, 201, 210, 216.	Citrus vulgaris, 267.
Catechu tannic acid, 216.	Cladophora, 221.
Cedar, 121.	Claviceps purpurea, 2.
— nut oil, 20.	Clupanodonic acid, 8.
Cellobiose, 53, 135.	Coagulation of proteins by heat, 287;
Celluloid, 143.	enzymes, 288.
Cellulose, 73; action of chemicals on,	Coco, 273, 278.
134, 135, 136; constitution of, 139;	- nut oil, 3, 29, 32, 33.
microchemistry, 144; occurrence,	Cocos butyracea, 3.
131; preparation of pure, 133; pro-	- nucifera, 3, 122, 132, 275.
perties of, 133, 134.	Codeine, 262.
Cell-wall formation and tannins, 220.	Co-enzymes, 354.
Centaurea cyanus, 249.	Coffee arabica, 122, 132. Coffee, 278.
Ceramium rubrum, 250.	Colchicine, 269.
Cera musæ, 43.	Cold and food reserves, 115, 305.
Cerasin, 126.	Collidine, 274.
Cerasus, 194.	Collodion, 142.
Ceratonia siliqua, 121.	Colloidal sodium chloride, 284.
Cereals, 121, 322, 323, 324, 387; leci-	Colloids, 291.
thins in, 44, 45.	Colour inheritance, 253, 256.
Cerotic acid, 43.	Colza oil, 2, 3, 4, 20, 22.
Ceryl alcohol, 1, 43.	Combretaceæ, 183.
Chara, 115.	Compositæ, 114, 180, 183, 396.
Chall devices 222	Compound celluloses, 137.
Chelidonium, 101.	Conifere 121 ISS 221
Chenopodiaceæ, 252.	Coniferæ, 121, 188, 321. Coniferin, 188.
Chenopodium Vulvaria, 49, 277.	Conifern, 188, 189.
Cherry gum, 57, 59, 126. Cherry-kernel oil, 19.	Coniine, 259, 262.
Chestnut, 121, 210, 216.	Conium maculatum, 266,

Conjugated proteins, 325. Diamino tri-hydroxydodecanic acid, 316. Copernicia cerifera, 43. Diamylose, 104. Copra, 3. Diarabinan tetragalactan-arabic acid, Cork formation and tannins, 220. Corydalis, 266. Diastase, 107, 108, 109, 118, 348, 368. Cotton, 132; fibre composition, 133; Dicranum, 117. seed, 74. Dictvota, 60. seed oil, 2, 3, 20, 32. Digallic acid, 193. Cranberry, 249. Digitalis purpurea, 119, 186. Crassulaceæ, 396. Digitogenin, 186, Crataegus Oxyacantha, 49, 275. Digitonin, 169, 186. Dioscorea japonica, 327. Diospyros Kaki, 219. Cress, 123. seed oil, 20. Croton oil, 20. Disaccharides, 53, 68. Cruciferæ, 187, 396. Divi-divi, 205, 207, 210, 216. Cucumis, germination, 40. Dracana australis, 120. — melo, 340. — rubra, 120. Cucurbita, 226, 315. Dragons blood, 204. maxima, 340. Drosera, 346, 372, 373. - Pepo, 279. Drosereceæ, 115, 176. Cucurbitaceæ, 183, 227. Drought, physiological, 115, 305. Currant, 128. Driers, 4. Cutch, 211, 218. Drying oils, 4, 20. Cuticle, 24. Dulcitol, 386. Cutin, microchemistry, 145. Dyer's broom, 246. Cutocellulose = Adipocellulose, 132, - sumach, 246. 138. Cyanidin, 249, 551. EARTH-NUT oil, 22. Cyanin, 248, 249. Echium vulgare, 195, 222. Cyanogenetic glucosides, 176; chem-Elaeis guineensis, 2, 3. istry, 178; microchemistry, 179; oc-Elaïdin test for fats, 21. Elaïoplasts, 35. currence, 176. Cyanophyceæ, 110. Elder, 121. Cycadaceæ, 127. Electrical energy and photosynthesis, Cynips aciculata, 195. 163. - gallæ, 212. Ellagic acid, 206. Cyperus esculentus, 2. Ellagitannic acid, 216. Cystine, 316, 329. Elodea, 115, 157, 161, 254, 257, 258. Cytase, 348. Emulsin, 75, 179, 181, 190, 348, 364, Cytisine, 268. 365. Cytisus Laburnum, 268. Emulsions, 14. Cytohydrolyst, 136. Emulsoids, 292, 293. Encrusting substances, 141. Dahlia, 111, 359. Entada scandens, 184. Enterokinase, 355. Dahlia variabilis, 321. Enzyme action of colloids, 297. Datiscetin, 247. Enzymes, action of light on, 351; and Datiscin, 247. Datisca cannabina, 2,7. antiseptics, 366; association, 347; Datura, 298. catalytic nature, 345; classification, - Stramonium, 281. 348; colloidal nature, 351; colour

DAHLIA, 111, 359.

Dahila variabilis, 321.

Datiscetin, 247.

Datiscin, 247.

Datisca cannabina, 247.

Datisra cannabina, 281.

Delphinidin, 249, 251.

Delphinin, 249, 269.

— consolida, 249, 251.

Delphinin, 249, 269.

— consolida, 249, 251.

Depletion of glucosides, 176.

Desmanthus, 191.

Dextrosa, 7, 103, 106, 110.

Dextrosae, 97.

Dextrose, v. Glucose.

Dhurin, 182.

Diamino acids, 316.

— nitrogen, 318.

348; colloidal nature, 351; colour inheritance and, 253; constitution, 349; mode of action, 352, 361; occurrence, 347; poisoning, 357; preperties, 350; specific nature, 353, 354. Equisactum, 236. Erepsin, 348. Ereptase = Erepsin, 371, 372. Ergot, 121, 276.

Ergotinine, 269. Ericaceæ, 396. Erythro-dextrin, 107, 110.

Erythrophyll, 221, 241.	Fruits, succulent, 128.
Erythroxylaceæ, 268.	Fucose, 53, 60.
Erythroxylon Coca, 267.	Fucoxanthin, 229, 241, 243.
Esparto grass, 132, 135.	Fucus, 127, 229.
Ethyl alcohol, manufacture, 389; oc-	Fundulus heteroclitus, 358.
currence of, in plants, 385; properties,	Fungi, 44, 107, 110, 156, 176, 194, 224,
388. Ethylchlorophyllide, 235.	280. Funkia, 35.
Eucalyptus, 74, 195, 201, 385.	Furfural, 57.
- occidentalis, 194.	Fusel oil, 375.
Eugenia caryophyllata, 385.	- user on, 3/3.
Eugenol, 189.	Cuch as
	GAGEA, 35.
Euonymus atropurpurca, 386.	Gaillardia, 35.
- europæus, 226, 385, 386.	Galactane, 97, 122.
Euphorbia, 97.	Galactose, 65; estimation, 80.
Euphorbiaceæ, 176, 367.	Galbanum resin, 202.
Eurotium, 334.	Galeopsis tetrahit, 234.
Euzanthone, 246.	Gallic acid, 195, 200, 201, 205, 214,
Evergreen plants, 116.	285.
	Galls, 195, 205, 210.
	Gallotannic acid, 193, 200, 213, 214;
FAGUS, 49.	constitution, 214; synthesis, experi-
- silvatica, 252, 275.	ments on, 210.
Fats, acetyl value, 33; acid value, 28;	Gall wasp, 212.
chemical properties, 12; colour re-	Cambier, 210, 211, 214.
	Gelatinization of colloids, 287.
actions, 22; constitution, 5; estima-	
tion, 24; extraction, 10; industrial	Gels, 284, 287, 294.
uses, 3; iodine value, 29; micro-	Genista tineteria, 246.
chemical reactions, 23; origin from	Genistin, 247.
carbohydrates, 36; origin from pro-	Gentiana lutea, 77, 247.
teins, 42; photosynthesis, 35; physical	Gentianose, 77.
properties, 11; physiology, 35; reac-	Gentisine, 247.
tions, 20, 21, 22; saponification, 13;	Geranium maculatum, 196.
saponification value, 29; solubilities,	— molle, 288.
12; translocation, 42.	Geraniol, 386.
Fatty acids, 6.	Germination, 315; of Allium, 40; of
Fehling's solution, 56, 77.	Arachis, 40; of Cannabis sativa, 40;
Fermentation, alcoholic, 374-385; of	of Cucumis, 40; of Lucerne, 68; of
_ amino acids, 339.	<i>Ricinus</i> , 37, 38, 41, 49; of sunflower,
Fermentative activity of yeast, 111.	36.
Ferns, 127, 194.	Glaucophyllin, 232, 233.
Festuca, 120.	Glaucoporphyrin, 233.
Fibrin, 356.	Gleicheniaceæ, 183.
Fibrinogen, 356.	Gliadins, 324.
Ficus, 194, 370.	Globulin, 323.
Fig, 370.	Globulose, 327.
Filices, 127, 194.	Glucoprotein, 325, 327.
Fisetin, 246, 251.	Glucosamine, 326.
Flagellates, 110.	Glucose, 60; estimation of-(a) gravi-
Flavellagic acid, 207.	metric, 88, 92; (b) polarimetric, 95;
Flavones, 244.	(c) volumetric, 79, 82, 83; preparation
Flavonol, 245.	and properties, 60; reactions, 61.
Flax, 132.	Glucosides, constitution, 170; micro-
Food stuffs, reserve, 115, 174.	chemistry, 179; occurrence of cyano-
Formaldehyde, detection, 152; estima-	genetic, 176; physiology, 173; varia-
tion, 148; formation from chloro-	tion in amount, 174.
phyll, 230; occurrence, in plants,	Glucosides and cultivation, 177.
153, 156, 157, 160, 163; polymeriza-	Glutamic acid, 315.
tion, 151, 155, 165; tests for, 153.	Glutelin, 324.
Formose, 151.	Glyceric aldehyde, 151.
Fraxinus Ornus, 386.	Glycerine, v. Glycerol
Fructose, v. Levulose.	Glycerol, 5, 13; reactions, 21.
26	
20	

404 IN.	DEA
Glycine, 315, 338.	Hexamethylene tetramine, 148.
Glycoamylin, 107.	Hexosephosphatase, 379.
Glycogen, 53, 97, 110; amount in	
yeast, 110; occurrence, 110; pre-	
paration, 112; properties, 113.	Histidine, 277, 316; occurrence, 321.
Glycogen-vacuoles, 111.	Histones, 322.
Glycogenase, 376.	Homogentisinic acid, 321, 360.
Glycollic aldehyde, 151.	Hops, 193, 194, 210.
Glycyl glycine, 319.	Hordenine, 276.
Glyoxaline, 277.	Hordenm sativum, 275, 279.
Glyoxylic acid, 338.	Hormones, 253.
Gold number, 291.	Hornbeam, amount of formaldehyde in,
Goodeniaceæ, 115.	153, 163.
Gooseberry, 128.	Horse-chestnut, 245, 321.
Gossypium herbaceum, 2, 3, 275.	Hübl's iodine value of fats, 29.
Gramineæ, 176, 183.	Humulus Lupulus, 193, 194, 210, 245,
Graminin, 120	275.
Granulose, 99, 100.	Hyacinthus, 115.
Grape seed oil, 32.	Hydnocarpic acid, 8.
- sugar, v. Glucose.	Hydrastine, 269.
Grapes, 199.	Hydrastinine, 269.
Gratiola officinalis, 301.	Hydrastis, 266.
Graviperception, 359.	Hydro-cellulose, 135.
Guanine, 278, 281.	Hydrocharis, 254.
Guarana, 278.	Hydrocyanic acid, 175, 181; estimation,
Gum arabic, 125.	200; occurrence, 176; physiological
— tragacanth, 126.	significance, 176, 177, 337; reactions,
Gums, 53.	179, 182.
Gun cotton, 142.	Hydrogel, 284.
Guttiferæ, 183.	Hydrogen peroxide, production of, in
Gymnosperms, 194, 266.	plant, 162.
Gynocardin, 173.	Hydrolysis, 12, 13.
**	Hydroquinone, 203.
Нæматін, 237, 325. Hæmatinic acid, 237; imide, 237.	Hydrosol, 284.
Hæmatinic acid, 237; imide, 237.	Hydroxy-phenylethylamine, 276.
Hamatococcus, 256, 257.	Hymenodictine, 269.
Hæmatoporphyrin, 237.	Hyoscine, 267.
Hæmoglobin, 236, 262, 325.	Hyoscyamine, 267.
Hæmolysis, 185.	Hypoxanthine, 278, 279.
Hæmolytic action of saponins, 185.	Hysteresis, 377.
Hæmopyrrole, 237.	Yearney and
Half-shadow polarimeter, 94.	IDAEIN, 249.
Halphen's reaction for cotton-seed oil,	Iller paraguayensis, 278.
Hamamelie ara	Illicium religiosum, 203.
Hamamelis, 210. Hedera, 66, 199, 301.	Imidazole, 277. Impatiens balsamifera, 132.
Helianthus, 121.	Indian yellow, 247.
- annuus, 226; germination, 37, 38, 39,	Indicane, 191.
v. Sunflower.	Indigo, 192.
- tuberosus, 114, 275.	Indigofera, 174, 177, 192.
Helicin, 172, 200.	- anil, 191.
Hemerocallis, 97.	- arrecta, 191.
Hemi-celluloses, 67, 120; occurrence,	- sumatrana, 191.
132.	- tinctoria, 191.
Hemp, 132.	Indoxyl, 192; occurrence, 194.
— seed oil, 20, 22, 32.	Inorganic ferments, 298.
Heracleum, 235.	Inosite, 387.
Hesperidin, 169, 204.	Insectivorous plants, 346.
Heuchera americana, 196.	Inulase, 118, 348.
Hevea braziliensis, 182.	Inulin, 53; biology, 116; occurrence,
Hexa-amylose, 104.	114; preparation, 117; properties,
Hexabromide test for drying oils, 22.	118.
, ,,	•

INDEX		
Inversion of cane sugar, 70, 71.	Levulose, 53, 64.	
Invert sugar, 64.	Lichenin, 97.	
Iodine value, determination, 29; table,	Liebermann's reaction, 307.	
32; of sunflower during germination,	Lignification, 137, 220.	
37∙	Lignified walls, 24.	
Iris germanica, 385.	Lignin, 132, 137; microchemistry, 144.	
- pseudacorus, 115, 120.	Lignocellulose, 131, 132, 136, 137.	
— Xiphium, 115.	Lignone, see Lignin.	
Irisin, 120.	Liliaceæ, 115, 121, 183.	
Isatis tinctoria, 191.	Lilium tegrinum, 98.	
Isobutyl acetic acid, 7.	Lime-tree, 121.	
Isochlorophyllin a and b, 232.	Linaria vulgaris, 17.	
Isoeugenol, 189.	Linolenic acid, 8, 20.	
Isohæmopyrrole, 237.	Linolic acid, 8, 20.	
Isolactose, 167.	Linum, 177, 178, 182.	
Isoleucine, 315, 321.	- usitatissimum, 2, 4.	
Isomaltose, 53, 73, 167.	Linseed oil, 2, 4, 20, 22, 32, 33.	
Isomerism, 54.	Lipase, 36, 37, 366; isolation from	
Isoquinoline, 266.	Ricinus, 367.	
Isovaleric acid, 7.	Lipoids, 10, 44; and respiration, 51;	
Ivy, 66, 199, 301.	physiology, 50.	
	Listera, 115.	
Language	Lobeliaceæ, 115.	
Japan wax, I.		
Juglans, 323, 387.	Loganiaceæ, 183, 268.	
Juncus communis, 98.	Lolium perenne, 288, 385.	
Jumperus Sabina, 385.	Lotoflavin, 183.	
Jute, 132.	Lotus arabicus, 177, 183.	
	Lotusin, 183.	
Kinase, 355, 356.	Lucerne, germination, 68.	
Kino, 201, 204, 210, 216	Lupeose, 53, 73.	
Kjeldahl method of estimating nitrogen,	Lupin, 79, 156, 321, 336, 359.	
	Lupinine, 268.	
340.	Lupinus, 128, 288, 323.	
Kola nut, 278.	- albus, 45, 321.	
	- luteus, 45, 121, 131, 268, 279.	
Labiatæ, 266, 396.	- niger, 268.	
Lacmoid, 202.	Lupulotannic acid, 193.	
Lacquer, 391.	Luteolin, 246, 251.	
Lactarius deliciosus, 2.	Lychnis chalcedonica, 184.	
Lactase, 348	Lycoperdon Bovista, 340.	
Lactic acid, 381.	Lycopin, 241.	
Lactosin, 77.		
Lamium, 161.	Lysine, 316; occurrence, 321.	
	M	
— album, 158.	MacLurin, 193, 203, 243.	
— maculatum, 235.	Madder, 68.	
Lanolin, 43.	Magnoliaceæ, 176, 183.	
Larix europæa, 76, 121.	Mahogany, 217.	
Lathyrus, 253.	Maize, 68, 324, 336.	
- sativus, 275.	— oil, 20.	
Laudanosine, 269.	Malpighiaceæ, 115.	
Lauraceæ, 176.	Malt, 108, 109, 373.	
Leather, 193.	Maltase, 73, 182, 348.	
Lecithin, 44, 230; formation, 41, 49;	Maltodextrin, 108.	
preparation from egg yoke, 45; phy-	Maltose, 53, 71; estimation, 81, 82,	
siological significance, 50; reactions,	83; occurrence, 71.	
46.	Malva parviflora, 128.	
Lecythidaceæ, 183.	Mandelonitrile glucoside, 172, 182.	
Leguminosæ, 44, 128, 176, 183, 265,	Mangrove, 195, 211.	
323, 396.	Manihot, 182.	
Lepidium, 127, 128.		
Leucine, 315, 338; occurrence, 321.	Manna, 74.	
Lencoplasts, 226.	Mannane, 97, 120.	
	Mannite, 36.	
Levulosanes, 97, 114.	Mannitol, 386.	

400	ט
Mannocellulose, 97.	
Mannose, 53, 68; estimation, 8o.	
Maple, 68.	
Marattiaceæ, 127.	
Maté, 207, 278.	
Medicago, 128.	- 1
- lupulina, 288.	
- sativa, 350.	
Melampyrum arvense, 301.	H
Melecitose, 53; occurrence, 76.	li
Melibiose, 53, 75, 167.	l j
Mellitis Melissophyllum, 235.	1
Melon seed oil, 20.	1
Menyanthes, 115.	1
Menyanthin, 181.	1
Mercurialis annua, 275.	
Mesocarpus, 194.	11
Mespilus germanica, 385.	1
Metaformaldehyde, 150.	-
Metapectic acid, 129.	1
Metapectin, 129.	1
Metapeptone, 327.	1
Metarabic acid, 125.]
Metellagic acid, 207.	1
Methyl alcohol, 385.	1
Methylchlorophyllide, 235.	
Methy ethyl maleinimide, 238.	
Methyl pentose, 59.	1
Micrococcus ureæ, 348.	1
Middle lamella, 130.	1
Millon's reagent, 308.	I
Mimosa, 211, 217.	
— catechu, 201.	1
— pudica, 195.	-
Mitochondria, 226.	-
Molasses, 69.	
Molisch's reagent for carbohydrates,	(
58	(
Monilia sitophila, 347.	(
Monocotyledons, 115, 118, 127, 266.	(
Monomolecular reaction, 361.	
Monosaccharides, 53, 57-68.	
Morin, 246, 251.	1.
Moringatannic acid, 193, 243.	0
Morphine, 262.	13
Morus tinctoria, 193, 246.	. (
Moulds, 347, 354.	
Mucilage, 53, 124, 127; occurrence, 127.	12
Mucilage sacs, 127.	1
Mucin, 325.	10
Mucor, 112.	
— stolonifera, 382.	(
Mulberry, 121.	0
Musa, 97, 219.	16
Muscari, 97.	10
— botryoides, 115.	1,
Muscarine, 48, 275.	1
Muscineæ, 127.	10
Mushroom, 373.	10
Mustard seed oil, 20.	1
Mutarotation, 170. Mycoderma aceti. 298, 349.	Č
my voucemu uven. 290, 349.	. (

Mycose = Trehalose, 53; occurrence, 73. Mycotrophic plants, 116. Mydaleine, 274. Myelin forms, 46. Myoporineæ, 115. Myosotis, 115. Myriophyllum, 127, 128. Myristica, 3. Myristic acid, 7. Myrobalans, 207, 210, 216. Myrosin, 188, 348. Myrtaceæ, 183, 268. Myrtillidin, 249. Myrtillin, 249. Myxomycetes, 110. NARCISSUS poeticus, 242. Neomeris, 115. - dumetosa, 128. Nepenthes, 346, 371, 373. Neurine, 48, 275. Nicotiana tabacum, 266. Nicotine, 266, 269, 270. Nitrates, reduction, 334. Nitrogen bases, 263; physiology, 280; estimation, 340; source of, for plant food, 333. Nitrogen content, 333. Nuclein, 326. Nucleoproteins, 325, 326. Nux vomica, 281. Oak, 210, 211, 217. galls, 210. - red, 208. Oat, 347. Oenidin, 249. Oenin, 249. Oenothera biennis, 256. Oil, amount in seeds, 2; biological significance, 2; transformation into starch, 2. and tannin, 222. Olae europæa, 2, 3, 36. Oleaceæ, 121, 183. Oleic acid, 7. Olein, 29. Olive oil, 2, 3, 19, 22, 29, 32, 33. Orange, 166, 347. seed oil, 20. Orchidaceæ, 128, 191, 266. Orchis, 115. - Morio, 67, 127. Orcinol, 58. Ornithine, 316. Ornithogalum, 35. — arabīcum, 227. Oryza sativa var. glutinosa, 101. Osazones, 56, 62. Osmosis, 251. Osmotic pressure of colloids, 284. Ouroparia catechu, 217.

	4-7
0 :1	/ Dontage and
Oxidases, 253, 262, 335, 349, 390-92, 394.	Peptase, 372.
Oxycellulose, 135.	Peptone, 327.
P. FONIA 26 102 202	Peroxidase, 253, 349; identification,
P.EONIA, 36, 123, 302. — officinalis, 132.	Persian manna, 76.
Pæony, 302.	Persimmon, 199.
Palm-kernel oil, 29, 32, 33.	Phæophyceæ, 260.
Palm oil, 2, 3, 4, 29.	Phæophytin, 233.
Palmaceæ, 121.	Phajus, 191.
Palmitic acid, 6, 7.	Phalaris arundinacea, 120.
Palmitin, 29.	Phanerogams, 127, 138.
Pangium edule, 175, 176, 177, 178.	Phase test, 234, 240.
Panicum, 182.	Phaseolunatase, 182.
Papain, 348.	Phaseolunatin, 182.
Papaveraceæ, 265.	Phaseolus, 194, 387.
Papaverine, 269.	- lunatus, 177, 181, 182.
Papaw, 348, 370.	- multiflorus, 196.
Papilionaceæ, 268.	- vulgaris, 372.
Paradextrane, 97, 114.	Phellonic acid, 138.
Paraformaldehyde, 150.	Phenyl alanine, 316, 321, 339.
Paragalactane, 97; occurrence, 122.	Phenyl ethyl alcohol, 339.
Paraisodextrane, 114.	Phlein, 97, 120.
Paraguay tea, 278.	Phleum pratense, 97, 120.
Paralysers, 356.	Phlobaphene, 208.
Paramannane, 97, 121.	Phloionic acid, 139.
Parapectic acid, 129.	Phloretin, 204.
Parapectin, 129.	Phloridzin, 173.
Parapectosic acid, 129.	Phloroglucinol, 201, 204.
Parchment paper, 135.	Phænix dactylifera, 122, 346.
Parenchyma, 188.	Phosphatides, 44.
Parsley, 273.	Phospholipins, 44.
Pasteur's solution, 375.	Phosphoproteins, 325.
Pastinaca sativa, 385.	Photosynthesis, 154, 230; extra cellu-
Pathological growths, 195.	lar, 157, 231; products of, 35,
Paullinia cupana, 278.	154, 175.
Pavy solution, 83.	— and energy, 160, 163.
Pea, 132, 322, 336.	Phycoerythrin, 248, 258; preparation,
- fat, 17.	259; reactions, 259.
— nut oil, 19.	Phycophæin, 248, 260.
Peach-kernel oil, 19.	Phyllin, 232.
Pear, 128.	Phytase, 238, 388. Phytelephas macrocarpa, 67.
— seed oil, 20. Pectase, 129, 288, 349, 355.	
Pectic acid, 129.	Phytin, 348, 388. Phytohæmatin, 262.
- bodies, 53, 128.	Phytorchlorin, 233.
— substances, 66, 288.	Phytosterol, 17, 43.
Pectin, 129.	Phytylchlorophyllide, 235.
Pectinase, 129, 130, 348.	Pinus, 44, 194, 196, 219, 226, 305.
Pectocellulose, 132, 138.	- cembra, 268.
Pectose, 129.	Pineapple, 68, 357, 370.
Pelargonidin, 249.	Pine bark, 210.
Pelargonin, 249.	Pine-seed oil, 32.
Pelargonium, 249.	Piper, 266.
Pelletierine, 268, 270.	Piperaceæ, 183.
Penicillium, 215, 334.	Piperidine, 264.
- glaucum, 121, 354.	Piperine, 266.
Pentahydroxybenzophenone, 193.	Pisangceryl alcohol, 43.
Pentosanes, 53.	— wax, 43.
Pentose, 53, 57; estimation, gravi-	Pistia, 194.
metric, 86; volumetric, 78.	Pisum, 288, 387.
Pepsin, 327, 348.	- sativum, 226, 266, 321, 323.

408 INDEX

Pittosporaceæ, 183.	Prunus Padus, 176.
Plasmatic membrane, 50.	Prussic acid, see Hydrocyanic acid.
Plasmolysis, 376.	Pseudo-cellulose, 132.
Platanaceæ, 176.	Pseudo-chloroplasts, 227.
Platanus, 236.	Pseudo-nuclein, 325.
Platinichlorides of bases, 270, 276; of	Pseudo-tannin, 193, 210.
lecithin, 46.	Psilotum, 35.
Pleurococcus, 334.	Pterocarpus, 201.
Plum-kernel oi!, 19.	- saxatilis, 128.
Polarimeter, 93, 94.	Ptomaines, 274.
Polemoniaceæ, 183.	Pulvini, 195, 218.
Pollen, 322.	Pumpkin, 321.
Polygalaceæ, 183.	— oil, 20.
Polygonum tinctorium, 191.	Punica granatum, 268.
Polymerization of aldehydes, 150.	Purine, 277.
Polypeptides, natural, 320, 327; syn-	Pyridine, 264, 266.
thetic, 319, 328.	Pyrocatechol = Pyrocatechin, 201, 202.
Polyporeæ, 194.	Pyrocatechol tannins, 210.
Polyporus betulinus, 114.	Pyrogallol, 204.
Polysaccharides, 97.	- tannins, 210, 211.
Pomegranate, 210, 216, 266, 268.	Pyrone, 247.
Pontederia cordata, 98.	Pyroxylin, 142.
Poppy-seed oil, 4, 20.	Pyrrole, 273.
Populin, 174.	Pyrrolidine, 267, 273.
Populus, 175.	Pyrroudine carboxylic acid = Proline,
- nigra, 245.	316.
- pyramidalis, 245.	Pyrroline, 273.
Porphyrin, 236.	Pyrrophyllin, 232, 233.
Portlandia grandiflora, 193.	Pyrroporphyrin, 233.
Potato, 185, 281, 392, 393.	Pyrus, 49, 194.
Precipitation of colloids, 304.	— Amygdalus, 181.
Precipitin, 330.	- Aucuparia, 65, 181, 235, 275, 338,
Primrose, 199.	386.
Primula sinensis, 253.	- cydonia, 181.
Primulaceæ, 183.	- Malus, 181, 246, 385.
Prochromogens, 261.	, , , , , , ,
Prolamin = gliadin, 324.	
Proline, 316, 328, 329; occurrence in	QUEBRACHO, 211.
plant, 321.	— colorada, 246.
Prosthetic group, 325.	Quercetin, 204, 242, 251.
Protamines, 322.	Quercitannic acid, 217.
Proteaceæ, 176, 183.	Quercitrin, 169, 245.
Protease, 370.	Quercus, 194, 195.
Protective power, 291.	— Cerris, 196.
Protein, animal and vegetable, 328:	coccinea, 195, 196.
colloidal properties, 308; composi-	— Ilex, 195.
tion, 317, 318; crystals, 301; de-	- infectoria, 212.
composition products, 281, 313;	- lusitanica, 195.
estimation, 343; extraction, 330;	- fedunculata, 195, 196.
grains, 301; hydrolysis, 317; micro-	- Prinus, 197.
chemistry, 308; occurrence, 301,	— sessiliflora, 195, 196.
329; optical activity, 304; origin of	- tinctorius, 169, 245.
fats from, 42; properties, 302; puri-	Quillaia, 184.
	Quinine, 268; preparation from cin-
333.	chona bark, 272.
Proteins and cold, 305.	Quince oil, 19.
Proteoses, 327.	Quinoline, 265.
Protoalkaloids, 273.	— alkaloids, 268.
Protocatechuic acid, 201.	Quinovin, 60.
Protoplasm, enzyme action, 367, 373.	Quinovite, 60.
Prunus Laurocerasus, 176, 178, 180.	Quinovose, 53, 60.
	Quino. 5.5, 55, 55,

	(1)
Panieu-seen oil 20	SACCHAROMYCES, 112, 375, 377, see
Radish-seed oil, 20. Radium emanation and photosynthesis,	also Yeast.
16c.	- Cerevisea, 110, 111; amount of
Raffinose, 53; occurrence, 74.	glycogen, 110.
Ranales, 176.	Saccharose v. Cane Sugar.
Rancidity of fats, 18.	Saccharum officinale, 68.
Ranunculaceæ, 176, 183, 265, 396.	Salep mucilage, 67, 121, 127.
Ranunculus bulbosus, 288.	Salicase, 190.
Rape-seed oil, 4, 20.	Salicin, 174, 175, 189; decomposition,
Raphidium, 334.	190; preparation, 189.
Ratanhia, 210.	Salicornia ramosissima, 252.
Ratanhia, 210. Reichert Meissl value determination,	Salicylic aldehyde, 190.
33; table, 33.	Saligenase, 190.
Rennin, 349.	Saligenin, 172, 189, 190.
Reseda Inteola, 246.	Salix, 175.
Reserve cellulose, 120; occurrence,	— viminalis, 189.
132; food stuffs, 174; of aquatic	
plants, 115.	Sambucus, 174, 236.
Resin, 196, 219.	— nigra, 176, 177, 182.
Resorcinol, 202.	Sambunigrin, 182.
Respiration, 51, 242, 258, 261, 334, 385,	Sapindaceæ, 176.
395. Reversible changes of colloids, 287.	Sapindus, 184. Sapogenin, 186.
	Saponaria, 184.
— reaction, 363. Reversion, 100.	— aiba, 183.
Rhamnaceæ, 183.	- officinalis, 107.
Rhamnetin, 246.	— rubra, 183.
Rhamnose, 53, 59.	Saponarine, 107.
Rhamnus cathartica, 216.	Saponification, 13, 14, 21.
- infectorius, 346.	- value, determination, 29; tables, 29.
- tinctoria, 246.	Saponins, 107, 183.
Rheum, 256.	Sapotaceæ, 176.
Rhinanthin, 173.	Sarracenia, 195, 218.
Rhizophora mangle, 195.	Saxifragaceæ, 176, 183.
Rhododendron, 194.	Saururaceæ, 183.
Rhodophyllin, 232.	Schizostylis, 115.
Rkus, 194, 391.	Schweitzer's reagent for cellulose, 134.
- coriaria, 195, 212.	Scilla, 154.
- cotinus, 246.	- maritima, 120.
- semialata, 212.	- nutans, 97, 98, 115.
— succedanea, 350.	— sibirica, 98, 115.
Ribes, 176. Rice, 324.	Scleroproteins, 324. Scrophulariaceæ, 396.
— oil, 19.	Scrophularia nodosa, 193.
Ricin, 274.	Scutellum, 345.
Ricinoleic acid, 8.	Seaweeds, 297.
Ricinus communis, 2, 36, 37, 38, 39, 44,	Secretory cells, 345, 346.
49, 274, 288, 301, 302, 367; germina-	Seed dispersal, 128.
tion, 37, 39, 41, 49; separation of	Selaginella, 225.
oil, 4.	Selective absorption, 297.
Robinia pseudacacia, 195, 218.	Semi-drying oils, 20.
Root, secretion of oxidase, 396.	Seminase, 225.
Rosaceæ, 176, 183, 266, 396.	Sepsine, 274.
Rosa gallica, 249.	Serum, albumin, 330.
Rosales 176.	— globulin, 330.
Rubiaceæ, 265, 268.	Sesame oil, 20, 22, 23, 32.
Rubia tinctoria, 193.	Silene vulgaris, 77.
Rumex hymenosepalus, 195.	Sinapis, 157.
Russula, 359, 392. Rutaceæ, 183.	— nigra, 187.
Rye, 121, 324.	Sinigrin, 187. Sinistrin, 120.
— oil, 20.	Sitosterol, 17.
011, =01	, Situaterol, 1/.

410 INI	DEA
Soap, detergent action, 14; manufacture, 5. — nuts, 184; solution, as a colloid, 14.	Sunflower, 37, 334, 336, see also Heli- anthus. — oil, 20.
— wort, 184.	Suspensoids, 292.
Soil as a colloid, 297.	Synanthrin, 97.
— and a tannin formation, 196.	Synthease, 381.
Soja bean oil, 20.	Syringa vulgaris, 386.
Soja hispanica, 122.	
- hispida, 132.	TAMARIX, 196.
Solanaceæ, 266, 267.	Tannase, 215.
Solanin, 185.	Tannic acid, 195.
Solanum dulcamara, 185.	Tannin, 193; biological significance,
- nigrum, 185.	223; distribution in plant, 218,
- tuberosum, 340, see also Potato.	219; economic use, 195; forma-
Solid spirit, 143.	tion in plant, 218, 222; occur-
Soluble starch, 102, 107.	rence, 194; microchemistry, 197; physiology, 217; variation in
Sorbic acid, 338.	amount, 196.
Sorbitol, 386. Sorbose, 53, 65.	— and anthocyanin, 254.
Sorghum, 177, 178, 183.	- cork formation, 220.
- saccharatum, 68.	Tannins, 193; chemistry, 200; classifica-
- vulgare, 182.	tion, 209; general properties, 193.
Soxhlet's method of extraction, 24.	— as glucosides, 208.
Sparganium, 115.	Taraxacum, 121.
Sparteine, 270.	Tartary soap, 184.
Spermaceti, 1, 43.	Taxicatin, 175.
Sphærocrystals, 118.	Taxus, 175, 236.
Spinach, 295.	Tea, 205, 210, 212, 243, 278, 279, 280. Terminalia Chebula, 194, 207.
Spinachia oleracea, 340. Spiræa Ulmaria, 190.	Thea sinensis, 246, 385.
Spiragyra, 157, 194, 195, 198, 220,	Thebaine, 269.
221.	Theobroma, fat, 2.
- crassa, 280.	— cacao, 128, 278.
Stachydrine, 267.	Theobromine, 277, 278, 281.
Stachyose, 53; occurrence, 76.	Thorn apple, 281.
Stachys silvatica, 235.	Thrombin, 349, 356.
— tuberifera, 76, 267.	Thrombogen, 356.
Starch, 115; estimation, 105; occur-	Thrombokinase, 356. Thymelæaceæ, 183.
rence, 97; preparation, 98; pro-	Tobacco, 273.
perties, 99; reactions, 104; transformation into oil, 2.	Tomato, 241.
- grains, chemical nature, 99; growth,	Toxic agents and enzyme action, 357.
101; physical nature, 101.	Tragacanthan-xylan-bassoric acid, 126.
Stearic acid, 7, 12.	Tragacanthose, 126.
Sterculia scaphigera, 127.	Tradescantia virginica, 98.
Stigmasterol, 69.	Translocation of fats, 42.
Stinging nettle, 238, 235, 240, 242.	Transpiration, 128, 257.
Stratiotes, 115.	Trehalose, 53, 73.
Straw, 132, 136.	Trianea, 194. Trifolianol, 386.
Strawberry, 68. Strelitzia, chloroplasts, 2.	Trifolium pratense, 279.
Strobilanthes, 191.	- repens, 279.
Strychnine, 268, 271.	Trigonellin, 266.
Strychnos Ignatii, 268.	Trigonellum fænum, 266.
- nux vomica, 268, 271, 281.	Trimethylamine, 49, 263, 273.
— toxifera, 268.	Triolein, 7.
Suberin, 139; microchemistry, 145.	Trioxymethylene, 150.
Sucrose, occurrence, 68; yield of, from	Trisaccharides, 53, 74-76. Trisetum, 120.
beet, 69.	Tristearin, 6.
Sugar, formation, 164. — and anthocyan, 254.	Triticin, 97, 120.
Sumach, 195, 205, 210, 212.	Triticum repens, 120.

Triticum sativum, 275, 278.

Tritona, 115.
Tropaelum, 155, 164.
— majus, 132.
Tropane, 267.
Tropistic movements, 359.
Trypsin, 327, 348, 355.
Trypsinogen, 355.
Tryptophane, occurrence in plant, 321; reaction, 373.
Tunicates, 131.
Tunicin, 131.
Turkish galls, 212.
Tyrosine, 307, 316, 359, 360; occurrence in plant, 321.

ULOTURIN Subilis, 280.
Ultra-violet light and photosynthesis, 160, 164.
Ulva, 2:20.
Umbelliferæ, 3g6.
Unsaponifiable residue of fats, 15.
Unsaturated compounds, 9.
Urea in plants, 279, 340.
Urease, 348.
Urica cid, 279.
Urine, 201.
Urita, 231, 236, 242.
Utricalaria, 127, 195, 218.

VACCINIUM Vitis Idea, 174, 202.

— Myrtillus, 385.
Valine, 315.
Valonia, 210, 216.
Vanilla, 35.

— planifolia, 189.
Vanillin, 189.
Varnish, 4.
Vaucheria, 2, 35,
Veratrine, 203.
Veratrum sabadilla, 203.
Verbascum Thapsus, 17.
Vicia Faba, 228, 123, 359.

Vicia sativa, 45, 275, 279, 321. Violaceæ, 115. Vigna sinensis, 323. Viscoid, 143. Viscose, 134, 143. Vitis, 387.

Walnut oil, 4, 20. Wash wood, 184. Water melon oil, 20.

Wax, chemical signifiance of term, 1; composition, 6, 44; function, 43; occurrence, 43, 230.

Weld, 246.
Wheat, 68, 121, 322, 324, 346.
— meal oil, 19.
Wijs' iodine value of fats, 31.
Willow, 174.
Wood gum, 57, 59.
Wound gun, 126.

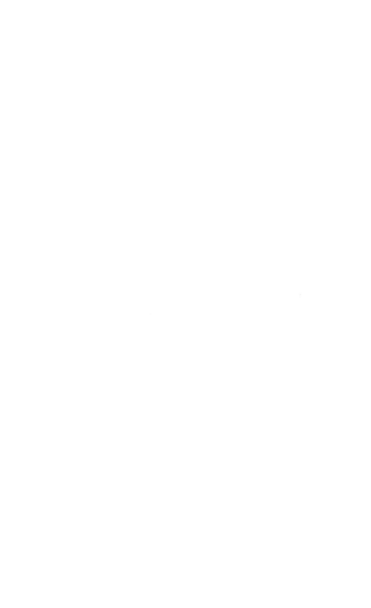
Xanthine, 277, 281. Xanthone, 244. Xanthophyll, 229, 241, 243, 262. Xanthoproteic reaction, 307, 308. Xeranthemum, 177. Xerophytes, 128. Xylose, 53, 59.

YEAST, 73, 108, 109, 121, 170, 278, 338, 373, 374; action on amino acids, 339; fermentative activity, 111; see also Saccharomyces.
Yellow wood, 246.

Yew, 199. Ylang Ylang, 385. Yucca, 115.

ZEA Mais, 177, 182, 345, 346, 385. Zygnema, 194. Zymase, 73, 349, 374-77, 383; quantitative estimation of activity, 383. Zymo exciters, 299.

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